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**SUMMARY REPORT INTERIM CLEANUP OF POLYCHLORINATED BIPHENYLS
(PCB) CONTAMINATED SOILS NEAR FORMER BUILDING 503**

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ERM-WEST

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INTERIM SUMMARY REPORT
INTERIM CLEANUP OF PCB CONTAMINATED SOILS
NEAR FORMER BUILDING 503

NAVAL STATION, TREASURE ISLAND
HUNTERS POINT ANNEX
SAN FRANCISCO, CALIFORNIA

DEPARTMENT OF THE NAVY
WESTERN DIVISION

NAVAL FACILITIES ENGINEERING COMMAND

SAN BRUNO, CALIFORNIA 94066-0727

A Report Prepared for

United States Navy
Western Division
Naval Facility Engineering Command
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SUMMARY REPORT
INTERIM CLEANUP OF PCB CONTAMINATED SOILS
NEAR FORMER BUILDING 503

NAVAL STATION, TREASURE ISLAND
HUNTERS POINT ANNEX
SAN FRANCISCO, CALIFORNIA

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Submitted March 3, 1989

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SECTION 1

INTRODUCTION

This summary report consolidates the information pertaining to the interim cleanup of PCB soils at a site near former Building 503 at Hunters Point Annex, (HPA), Treasure Island Naval Station (Figure 1-1). It includes the discovery of the PCBs, the subsequent investigations and studies to define the extent of the PCB contaminated soils, and the resulting cleanup and soil verification reports.

The interim cleanup took approximately two years - from discovery in the fall of 1986 to final cleanup and finishing of the site in the summer of 1988. Additionally, the cleanup involved the close working relationships of the Navy, the Department of Health Services, the Regional Water Quality Control Board, consultants, and remediation contractors. The technical cleanup work was directed and coordinated by the Western Division Naval Facilities Engineering Command (WESTDIV).

The work at the PCB site focused on the interim removal of PCB contaminated surface soils. The cleanup involved excavating and disposing of PCB soils at a Class I landfill. The cleanup goal for the site was defined as 25 mg/kg of residual PCBs. Appropriate long-term remedial actions for the site will be implemented, as required, after the completion of a Remedial Investigation/Feasibility Study (RI/FS) currently in progress at HPA.

This summary report is divided into three remaining sections. Section 2 provides the background from discovery to development of a workplan for cleanup of the site. Section 3 describes the interim cleanup work including methodology, soil disposal, field sampling and field analysis, final confirmation sampling, the results of the verification sampling program, and site finishing and future work. Section 4 is a reference listing. Figures and Tables referenced in the text of the report are consolidated in Appendix A.

SECTION 2

DISCOVERY AND SITE INVESTIGATIONS

This section describes the background and the investigations to identify the extent of the PCB site contamination. The PCB site, as shown in Figure 2-1, is located at the Hunters Point Annex, Treasure Island Naval Station on the west side of the Hussey Street and near the southeast corner of the new Shore Intermediate Maintenance Activity (SIMA) building. The studies and subsequent reports prepared during the discovery phase of the project are summarized in this section.

DISCOVERY

The first indication of PCB contamination at the site was in September of 1986 when the Naval Public Works Department (PWD) was excavating soil to locate and to repair an underground utility. The location of the excavation was between the former building 503 and Sanitary Sewer (SS) Pump Station No. 1 on the western edge of Hussey Street. The location is identified as sample point 1 on Figure 2-2. After finding stained soil and odor in the excavation, the contractors notified the Naval Public Works Department (PWD). After soil samples were taken by PWD and field analysis confirmed PCB contamination, the Western Division of the Naval Facilities Engineering Command (WESTDIV) was consulted for further advice and action.

ERM-West was commissioned to assist WESTDIV to further investigate and perform the following tasks: 1) confirm the PWD results; 2) develop and complete a general survey of the SIMA site for other possible PCB soils; and 3) develop and complete a survey of ground water to assess potential contamination in the area.

Each of the three investigations resulted in reports that are summarized in the following paragraphs.

CONFIRMING THE DISCOVERY

On September 12, 1986 ERM-West collected four samples in the general area of the PWD excavation and requested testing for PCBs by a certified hazardous waste analytical laboratory (ANLAB Analytical Services, Sacramento, California). Soil samples confirmed the presence of PCBs in the soils but in concentrations that would not classify the soils as hazardous by either the State of California or the Environmental Protection Agency (results ranged from less than 0.001 ug/l to 2.3 ug/l in standing water in the excavation, to 0.02 mg/kg in soil). Results of the findings were presented to the Navy in a report dated September, 1986.¹

General PCB Site Survey

After confirming PCBs in the soil the recommendation was to survey the SIMA site more extensively. The extent of the site survey performed on October 15 - 16, 1986 is shown in Figure 2-2.² Sixty borings were drilled to a depth of approximately five feet. Additionally, two borings were drilled to depths of greater than six feet to obtain groundwater samples. Soil samples were generally composited for each boring, and then analyzed for PCBs by a two step procedure: 1) a field test kit procedure to identify potentially high concentrations of PCBs, followed by 2) a certified hazardous waste laboratory analysis by gas chromatography (GC) on selected high concentration samples identified by the field test kit.

All soil samples were initially subjected to field testing in an on-site laboratory constructed and operated by ERM-West

personnel. The field test kit was manufactured by McGraw Edison Company and has been demonstrated to be an accurate in-field tool to assist in the identification of PCBs. (For further discussions for limitations and constraints of the field test kit see reference 2). Additionally, two soil samples were submitted to a certified laboratory (ANLAB) for purgable halocarbons (EPA method 8010) analysis to determine if other contaminants had leached into the soil near a separator and sump adjacent to the transformer pad.

The results of the survey confirmed that PCB soil contamination appeared to have spread from the old transformer pad, located in the southeast corner of former building 503, toward Hussey Street. In Figure 2-2, the sample points are identified by a number within a circle. Four of the sixty samples between the transformer pad and Hussey Street had high PCB soil concentrations ranging from 26 mg/kg to 910 mg/kg of Aroclor 1260. These sample locations are identified in dark blue on Figure 2-2 and are identified as sample numbers 2, 2A, 2Q, and 2R. A summary of the field kit analysis and the certified laboratory results are presented in Appendix B for reference.

No detectable contamination was reported for the remainder of the samples collected in the survey. No detectable purgable halocarbon were found in the soil. The report recommended development of a work plan to cleanup the site.

Ground Water Surveys

Ground water studies were performed in January and September, 1987. In January, 1987, an investigation³ of soil, sediment, surface water, and groundwater was conducted to determine if PCBs were present above action levels. The field work included logging and sampling of borings; installation, development, and sampling

of seven wells; water level measurements; and sampling of sediment and water from a catch basin.

The location of the seven monitoring wells, identified as W-1 through W-7, are shown in Figure 2-3. The January, 1987 investigation indicated that groundwater gradient was non-uniform and appeared to flow in an easterly direction. Low concentrations of PCBs (0.004 mg/l) were detected in one of seven wells (W-1). Well W-1 is located at the transformer pad and, based on the water samples taken during the study, there was no indication that groundwater was spreading the PCBs.

In September, 1987, ERM-West prepared a report for a ground water monitoring plan^{3A}. This study combined the data collected from the January 1987 investigation with new data collected from the existing seven monitoring wells and the drilling of four additional borings (B8 through B11 shown in Figure 2-3). The report summarized the site's soil lithology, groundwater gradient, tidal influences, aquifer characteristics, and the occurrences of PCB at various soil depths and within the groundwater. In Appendix C are tables of data summarizing the results of PCB analysis on subsurface soil and groundwater. Additionally, geologic cross sections of the site are presented in Appendix C.

Trace but detectable concentrations of PCBs in groundwater were reported in two of five wells at the site. These two wells (W-1 and W-2) are within the area where surface and subsurface PCBs had been detected and were within the "footprint" of the PCB cleanup site. The levels of PCB detected ranged from 0.0007 mg/l to 0.004 mg/l.

PCBs were found in the subsurface soils at depths as great as 27 feet. The concentrations at depths greater than 20 feet were typically trace concentrations, less than 0.1 mg/kg. The highest

concentrations, however, were detected in well W-1 located at the former transformer site (highest values ranged from 24 mg/kg to 32 mg/kg). With two of the highest concentrations (28-32 mg/kg) occurring within 5.5 feet of ground surface. This surface contamination was removed with the soil cleanup activities that occurred at the site.

Four wells were subsequently sealed and destroyed (Wells W-1, W-2, W-3, and W-5) by new construction (SIMA Building) and soil excavation during the PCB interim remediation work. Three of the wells (W-4, W-6, and W-7) remained as ground water monitoring wells.

The groundwater monitoring plan recommended the installation of two new monitoring wells. With the three existing wells at the site, the monitoring well network proposed would have incorporate data from five wells. The installation of the two additional wells was not implemented by the Navy. After a meeting between the Navy, Regional Water Quality Control Board, and Department of Health Services, an agreement was made that the information in the groundwater monitoring report would be included in the RI/FS study that is presently in progress. The decision to place monitoring wells at specific locations within the Hunters Point Annex, as well as this PCB site, would be determined as part of the overall RI/FS study.

SECTION 3

INTERIM CLEANUP SUMMARY

This section summarizes the interim cleanup work at the PCB site near former Building 503, Hunters Point Annex, Treasure Island Naval Station. The cleanup of PCB contaminated surface soils comprised site management and contractor coordination, development and preparation of a Health and Safety Plan, determination of the spill site boundary, cleanup by excavation and removal of soils to a Class I hazardous waste landfill, and verification that the cleanup goal for surface soils had been achieved by the soil excavation work.

The cleanup work was performed in three phases. A report^{4,5,6} was prepared at the completion of each work phase to summarize the excavation activities, the soil sampling results, and the final verification sampling results. The Navy would authorize the cleanup activities to proceed onto the next cleanup area of the site after review and comment by the regulatory agencies of the results and recommendations of the previous work phase.

The results of the cleanup work are presented in the following paragraphs. The total yards of contaminated soil removed from the site and disposed in a Class I landfill was approximately 1500 cubic yards. Landfills used as disposal sites were Chemical Waste Management in Kettleman City, California and United States Pollution Control Inc. at Grassy Mountain, Utah.

PROJECT ORGANIZATION AND RESPONSIBILITY

Four groups were directly involved in the cleanup of the site: the Navy, an excavation and disposal contractor, an environmental engineering consultant, and an analytical laboratory. The PCB cleanup project was directed by the Western Division Naval Facilities Engineering Command (WESTDIV), San Bruno, California. WESTDIV coordinated the cleanup work using contractors and consultants, communicated with the regulatory agencies, inspected the site, and tracked work progress.

The cleanup contractor was American Environmental Management Corp., Oakland, California. The contractor was responsible for excavation and removal of soil from the PCB site, site containment, and health and safety of American Environmental personnel.

Sampling of the site was performed by ERM-WEST, Walnut Creek, California. ERM-WEST was responsible for sampling the site and reporting the PCB results to the Navy; setting-up the verification sampling grid; overseeing health and safety of ERM-West personnel; and reporting sampling results.

Central Coast Analytical Services, San Luis Obispo, California, was the laboratory used to perform the PCB analytical procedures. Central Coast Analytical Services is a certified hazardous waste laboratory and was used during all three cleanup phases.

HEALTH AND SAFETY PLAN

ERM-West prepared a Health and Safety Plan specifically tailored to the work to be performed by ERM-West personnel.⁷ Since the contractor, American Environmental Management Corp, was involved in the physical removal of the soil, a separate Health and Safety Plan was prepared by American Environmental for their personnel.

CLEANUP GOAL

In a February 18, 1987 letter⁸ from the Department of Health Services (DHS) to the Navy, the DHS concurred with EPA's recommended PCB cleanup level of 25 ppm. This cleanup goal for surface soils had a condition that any areas having a residual PCB concentration greater than background levels would be covered over by concrete, asphalt, or a minimum of 10 inches of clean soil.

SOIL EXCAVATION

The extent of the cleanup area is shown in Figure 3-1. The basic shape of the entire cleanup area was defined in Phase 1 when soil excavation and disposal work was performed in all three areas of the site.

In order to optimize resources and to insure adequate removal and verification of contaminated soils, the excavation area was divided into three subareas. The subdividing of the site provided several advantages: 1) the large cleanup area was divided into manageable areas, 2) dewatering of the ground water was lessened by reducing the size of the cleanup areas, 3) in-field soil

sampling could be managed more effectively, and 4) final verification sampling grids were tighter and more encompassing than if the area was treated as one large site.

VERIFICATION SAMPLING PLAN

The determination of the PCB contamination boundary and the horizontal and vertical extent of soil contamination was based on the results of field samples analyzed by the McGraw Edison PCB test-kit. Use of the test kit as a screening device to direct the field activities in all three phases of work minimized the analytical costs and expedited the clean-up activities.

The decision to implement a formal verification sampling plan in each of the three phases at the work site was based on initial soil screening by the field PCB kit. The correlation between field sample results and certified laboratory reports has been confirmed and previously discussed in both the Interim Report⁴ and addendum.^{4A} Comparison of split samples analyzed concurrently by the field test kit and a certified analytical laboratory consistently indicated that the field kit conservatively indicated PCB concentrations higher than the certified laboratory results.

The following paragraphs summarize the verification sampling protocol used in the three cleanup areas. Detailed laboratory reports for the verification sample analysis by area are presented in Appendix D.

Sample Grid Layout Procedures

The layout of the verification sampling grid for each the three cleanup areas is shown on Figure 3-2 (Enclosed in pocket at

end of report). The location of sampling points is based on USEPA protocol for verification sampling⁹.

The sample points for each area were established as follows: the longest length of each spill area was determined (L1); at the midpoint of L1, a perpendicular line (L2) was established across the excavation; a line connected from the midpoint of L2 (center of the sample grid) to the endpoint of L1 was established as the radius of the excavation (this radius defines the extent of the spill area by the EPA protocol). Once this midpoint is established, verification sampling points are determined by creating a hexagonal grid based on equilateral triangles. The grid starts at the center of the of the sampling grid and radiates outward.

Once the radius of the sampling area was determined, the spacing of the hexagonal sample points was determined by multiplying the radius by 0.3 (Table 2⁹). To summarize, the sampling grid layouts for each area were as follows:

<u>Cleanup Area</u>	<u>Sample Radius, feet</u>	<u>Sample spacing, feet</u>	<u>Number of Samples</u>
1	50	15.0	37
2	37.5	11.3	50
3	26	7.8	56

In each phase of the clean-up work, additional grid points were identified on the outside periphery of the excavation to confirm that the extent of the surface PCB contamination had been accurately identified. Therefore, in each clean-up area, the total number of grid points sampled and analyzed equaled or exceeded the sample size referenced in Table 4⁹.

As indicated on Figure 3-2, nine of the sample points were triangulated and located relative to either the existing pumping station or existing power poles. These nine points allow re-establishment of the sampling grid at a future time.

Soil Sampling and Laboratory Results

Depths of the excavation ranged from approximately 1 to 10 feet, 1 to 6 feet, and 1 to 8 feet below the existing grade in areas 1, 2, and 3, respectively. Groundwater in the area was typically 2 to 4 feet below grade.

Different sampling protocols were used to sample the interior and outside periphery of the excavations. For the interior of the excavation, groundwater seepage in the excavation was pumped into an on-site holding tank, sampled, and analyzed for PCBs. Laboratory results did not detect PCBs and the groundwater was discharged to the pumping station. At each grid point, a 6-ounce surface soil sampled was taken with a wooden disposable tongue depressor and placed in a glass jar; depth of sample was approximately 2 centimeters. The Teflon-capped sample glass jars were placed on ice for transport to the laboratory.

The periphery of the site, outside the excavation, was sampled using a hollow stem drill rig. The drill rig was used to break through the asphalt paving and to push brass tubes for retrieval of undisturbed samples at depths varying between 1.33 feet and 3.0 feet. The depth of sample was previously discussed with the DHS before drilling and follows the procedures established during the Area 1, 2, and 3 sampling programs.^{4,5,6} Samples inaccessible to the drill rig were hand augered. After sample retrieval, sample tubes were covered with Teflon, capped, placed on ice, and transported to the laboratory.

In the following paragraphs, a summary of each area's verification sampling history is presented. Figure 3-2 depicts the sample locations and Tables 3-1, 3-2, and 3-3 summarize the final sampling results for each of the areas.

Area 1. On April 15, 1987, 21 samples from excavated Area 1 were composited in five groups. Three of the five composites had arithmetic averages greater than the cleanup level (25 mg/kg). Thus discrete samples were analyzed from these composite groups. Since the discretes for 75 percent of samples from the composite group of A1, A2, B1, B2, exceeded the cleanup goal, this area was included for cleanup in the Area 2 excavation work.

Of the remaining discrete samples only one sample, G4, exceeded the cleanup level. This area was further excavated following discussion between the Navy and DHS and was resampled on May 1, 1987. Additionally, sixteen samples, designated X1 through X16, were sampled from the periphery of the excavation on May 18, 1987. As presented in Table 3-1, the final verification sampling results indicated that no final verification sample exceeded the cleanup goal of 25 mg/kg.

Area 2. In area 2, the grid points were initially sampled on September 3 and 4, 1987. Areas where soil contamination exceeded the cleanup criteria of 25 mg/kg PCBs were further excavated and were resampled on October 16, 1987. Seven and two randomly split samples were shared with the DHS on September 3-4, 1987 and October 16, 1987, respectively. These samples were taken to the DHS laboratory for sample preparation, splitting, and subsequent forwarding of half of each sample to Central Coast Analytical Services for PCB analysis. As presented in Table 3-2, the final verification sampling results indicated that no sample exceeded the cleanup goal of 25 mg/kg. The split samples analyzed by the DHS were consistent with the results of Table 3-2.

Area 3. In area 3, the interior and the periphery of excavation were sampled on August 5 and August 8, 1988, respectively. No sample splitting was performed following discussions with the DHS. All samples collected during this work were reported below the cleanup goal of 25 mg/kg as indicated by the results summarized in Table 3-3.

Sample Tracking and Laboratory Reporting

To track samples from the field to the laboratory, chain of custody documentation was prepared and sealed with the field-to-laboratory transport container. Chain of custody forms prepared and used during the sampling program are presented in Appendix E.

ANALYTICAL PROCEDURES

Once the soil samples were received at the Central Coast Analytical Services laboratory the samples were prepared, extracted, and analyzed for PCBs. Sample preparation and extraction generally followed EPA method 3550 - Sonication Extraction. However, instead of a 1mm X 1mm sieve required by the EPA method, Central Coast used a #10 sieve (2mm X 2mm) to be consistent with sample preparation required by California Administrative Code, Title 22, Division 4, Chapter 30, Section 66700 (c) (1). This modification to the EPA method was implemented at the request of the DHS.

The samples received at the laboratory were air dried (Phases 2 and 3 only) before analysis for PCBs per DHS recommendations. This was done after determination of percent moisture in the original sample. A 20 gram wet weight sample was used and mixed with Na₂SO₄ then sonicated. The extraction solvent was 1:1

hexane:acetone (B & J) chromatography grade. The extract sample was filtered and reduced in volume using a Kuderna-Danish concentrator. PCBs, if present in the sample extract, were identified using EPA method 8080 - Organochlorine Pesticides and PCBs.

Central Coast's "Quality Assurance/Quality Control Procedure" as summarized in Appendix F, required blanks, duplicates, and spikes to be analyzed once per batch, once per matrix type or once per 20 samples, whichever is more frequent. For this project, however, duplicates were run once per batch, once per matrix type, or once per 10 samples analyzed, whichever was more frequent.

Background and Blank Samples

As part of the quality assurance program, a soil background (BL-1) and blank (BL-2) were collected during Area 2 verification sampling. Trace levels of PCBs, 0.17 mg/kg and 0.5 mg/kg were detected in both soil samples BL-1 and BL-2, respectively. Both of the occurrences were a result of factors not directly influencing the verification sample data. Compared to the PCB concentrations found in the actual verification samples, those found in samples BL-1 and BL-2 are not significant.

INTERIM CLEANUP APPROVAL, SITE FINISHING, AND FUTURE SITE WORK

Laboratory analysis of the soil samples obtained in Phases 1, 2, and 3 of the Interim Cleanup of the site using EPA Verification Sampling Protocol indicated that the interim cleanup goal of 25 ppm or less had been achieved for all three areas. Each phase of the cleanup work received DHS review and comment prior to site backfilling.

Each excavated area was backfilled with acceptable engineered fill and compacted to minimize soil settlement. The entire cleanup area was topped with asphalt paving to prevent surface water infiltration and surface erosion.

The PCB removal area will be further investigated in the Hunters Point Annex Remedial Investigation/Feasibility Study (RI/FS) to further characterize remaining low-level PCB contamination at the site. Upon completion of the RI/FS, long-term remedial measures to protect the public health and environment will be implemented as appropriate.

SECTION 4

REFERENCES

The following references were developed or used during the cleanup work at the PCB site near former building 503, Hunters Point Annex, Treasure Island Naval Station.

1. ERM-West, September 1986, Hunters Point Naval Shipyard, PCB Investigation
2. ERM-West, November 1986, Hunters Point Naval Shipyard, Preliminary Investigation of Possible PCB Spill
3. ERM-West, January 1987, Investigation of PCBs in Soil and Groundwater at the Hunters Point Site
- 3A. ERM-West, September 1987, Groundwater Monitoring Plan, Hunters Point Naval Shipyard, PCB Spill Site, Near Former Building 503
4. ERM-West, April 1987, PCB Verification Sampling Results, Interim Report (Area 1)
- 4A. Addendum to the "PCB Verification Sampling Results, Interim Report", Attachment to a letter from the U.S. Navy to the Department of Health Services, June 25, 1987.
5. ERM-West, December 1987, PCB Verification Sampling Results, Area 2
6. ERM-West, July 1988, PCB Verification Sampling Plan, Area 3

7. ERM-West, February 1987, Hunters Point - PCB Cleanup Work:
Health and Safety Plan
8. February 18, 1987, DHS letter to Alex Dong, Western Division-
Naval Facilities Engineering Command
9. USEPA, August 1985, Verification of PCB Spill Cleanup by
Sampling and Analysis, Interim Report No. 2

APPENDIX A
FIGURES AND TABLES

APPENDIX A
FIGURES and TABLES

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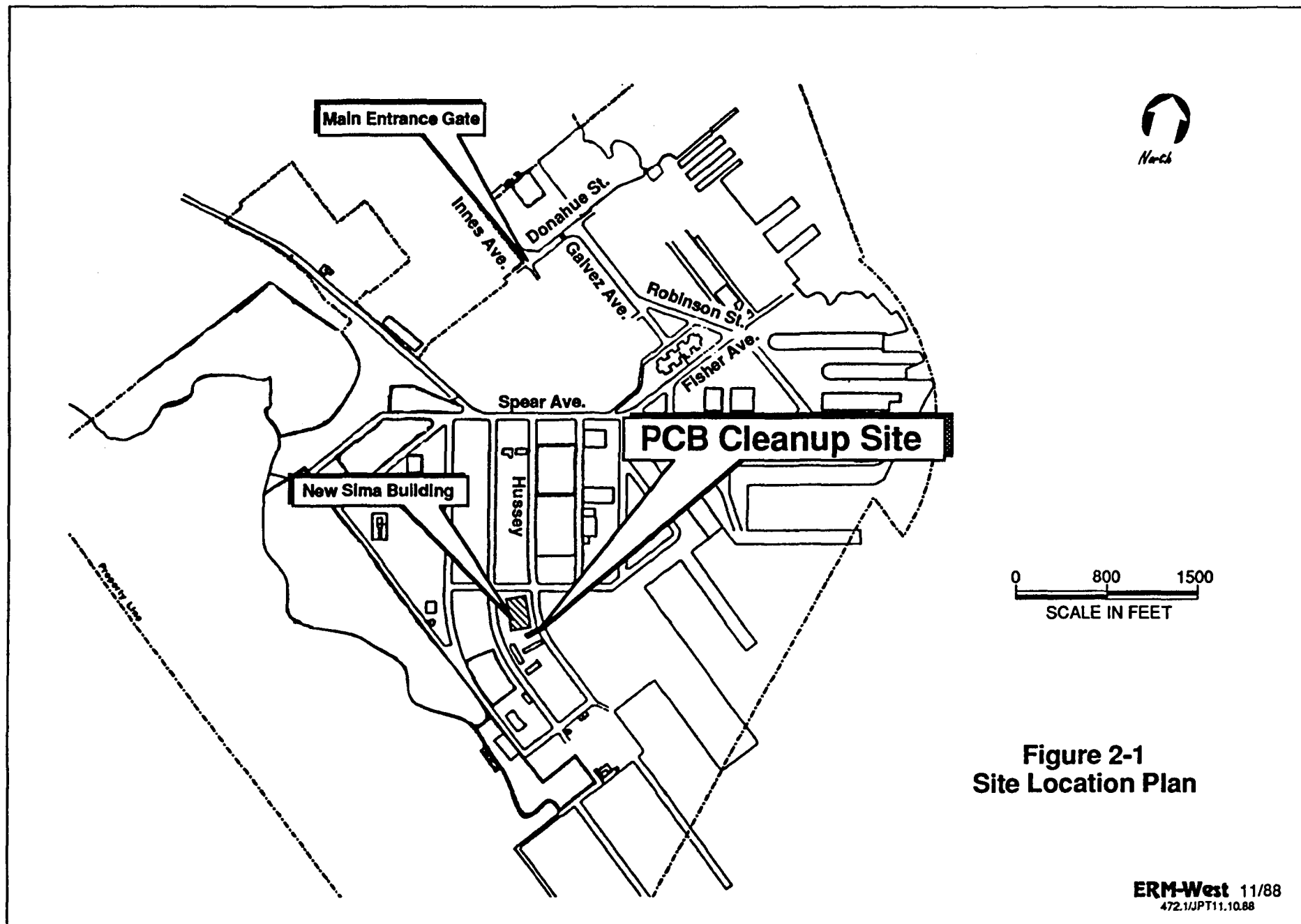


Figure 2-1
Site Location Plan



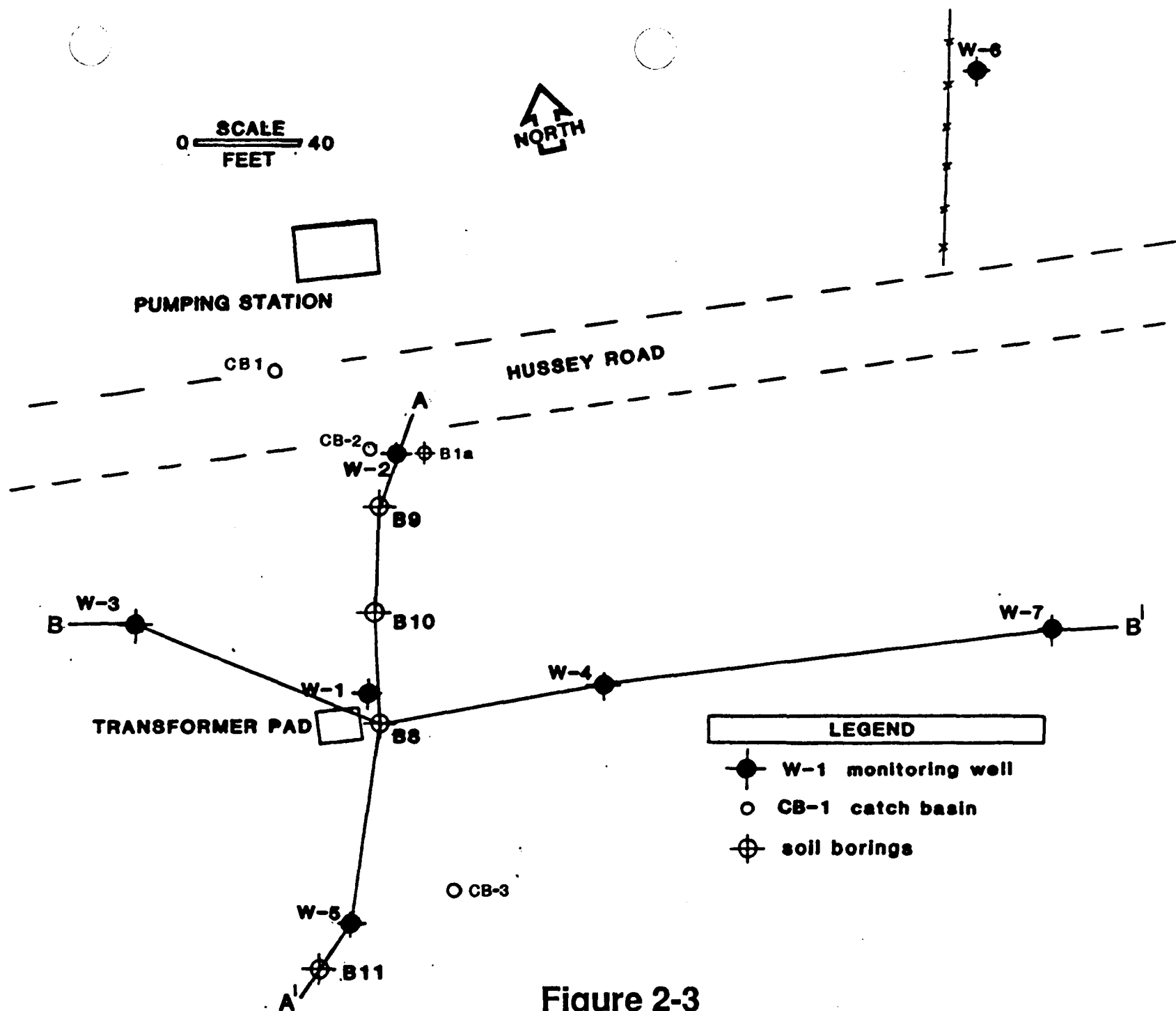


Figure 2-3
Locations of Borings, Wells and Geologic Cross Sections

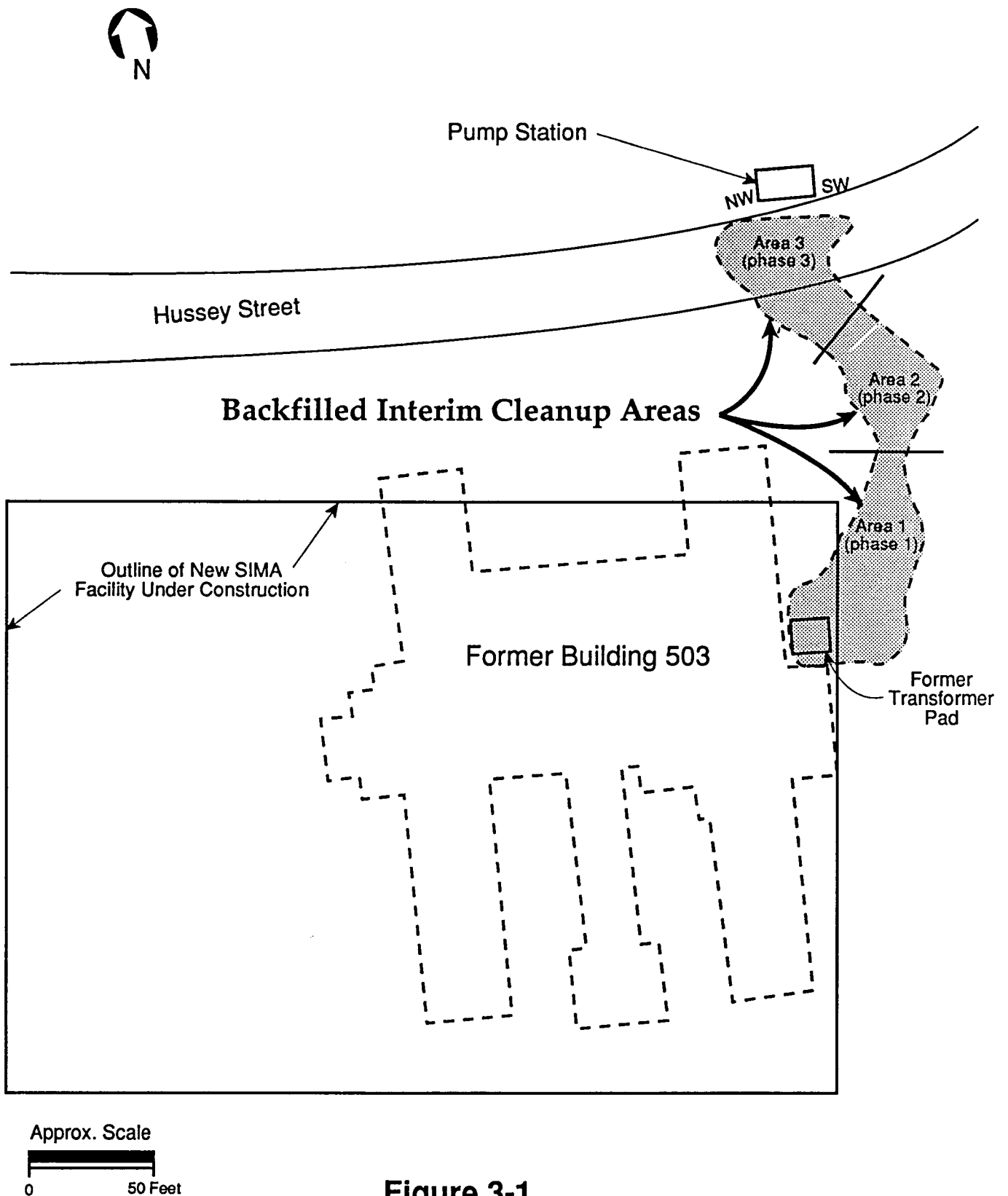


Figure 3-1
PCB Site-Work Areas

TABLE 3-1

HUNTERS POINT ANNEX
TREASURE ISLAND NAVAL STATION

VERIFICATION SOIL SAMPLING RESULTS
FROM CENTRAL COAST ANALYTICAL SERVICES
FOR AREA 1

April 15 and May 18, 1987			
Sample Description	Percent Moisture	PCBs ¹ D.L. ²	mg/kg Concentration ³
F2	NA	0.1	3.5
F3	NA	0.1	1.0
G1	NA	0.1	4.2
G2	NA	0.1	0.4
H1	NA	0.1	2.0
H2	NA	0.1	0.4
H3	NA	0.1	3.6
H3 (Dup)	NA	0.1	6.6
G4 (5/1/87)	NA	0.01	0.41
X1	NA	0.01	0.30
X2	NA	0.01	0.49
X3	NA	0.01	0.03
X4	NA	0.01	0.08
X5	NA	0.01	0.02
X6	NA	0.01	0.02
X7	NA	0.01	0.04
X8	NA	0.01	<0.01
X9	NA	0.01	0.04
X10	NA	0.01	0.10
X11	NA	0.01	0.04
X12	NA	0.01	0.05
X12 (Dup)	NA	0.01	0.05
X13	NA	0.01	0.25
X14	NA	0.01	0.10
X15	NA	0.01	0.08
X16	NA	0.01	0.16

Notes:

1. Only PCBs detected were Aroclor 1260.
2. D.L. is the detection limit.
3. Reported concentration of PCBs by the laboratory analysis.
4. NA - Not analyzed.

TABLE 3-2

HUNTERS POINT ANNEX
TREASURE ISLAND NAVAL STATION

VERIFICATION SOIL SAMPLING RESULTS
FROM CENTRAL COAST ANALYTICAL SERVICES
FOR AREA 2

September 3-4 and October 16, 1987

Sample Description	Percent Moisture	PCBs ¹ , D.L. ²	mg/kg Concentration ³
2A-1	6.0	0.2	<0.2
2A-2	3.2	0.02	5.0
2A-3	7.1	0.02	2.2
2A-4	2.7	0.02	0.61
2B-1	NA	0.02	<0.02
2B-2	16	0.02	16
2B-3	7.2	0.02	0.44
2B-3 (Dup)	7.1	0.02	--
2B-4	5.1	0.02	1.0
2B-5	3.8	0.02	0.60
2B-6	NA	0.02	<0.02
2C-1	NA	0.02	0.04
2C-2	30	0.02	25
2C-3	12	0.02	3.2
2C-4	25	0.02	15
2C-5	NA	0.1	1.3
2C-5A	15	0.05	2.4
2C-5A (Dup)	15	0.05	2.7
2C-5A (Tripl)	15	0.05	2.9
2C-5B	21	0.05	11
2C-5C	19	0.05	1.5
2C-6	4.7	0.02	0.65
2C-7	NA	0.02	<0.02
2D-0	NA	0.02	<0.02
2D-1	30	0.02	3.5
2D-2	24	0.05	16
2D-3	31	0.02	4.3
2D-4	21	0.02	2.2
2D-5	1.2	0.02	2.2
2D-6	NA	0.02	<0.02

TABLE 3-2 (continued)
 VERIFICATION SOIL SAMPLING RESULTS
 FOR AREA 2

September 3-4 and October 16, 1987

Sample Description	Moisture Percent	PCBs ¹ , D.L. ²	mg/kg Concentration ³
2E-1	5.8	0.2	<0.2
2E-1 (Dup)	5.8	0.2	<0.2
2E-2	18	0.05	3.0
2E-2 (Dup)	18	0.05	3.0
2E-2A	5	0.05	2.2
2E-3	NA	0.05	7.9
2E-3A	20	0.05	15
2E-3B	16	0.05	0.79
2E-4	19	0.05	8.9
2E-4 (Dup)	22	0.02	25
2E-4 (Spike)	19	0.05	14
2E-5	20	0.02	4.6
2E-5 (Dup)	20	0.02	4.2
2E-6	17	0.02	0.6
2E-7	NA	0.02	<0.02
2F-1	NA	0.02	<0.02
2F-2	25	0.02	3.4
2F-3	30	0.02	9.6
2F-4	17	0.02	8.4
2F-5	23	0.02	0.73
2F-6	1.6	0.02	0.6
2F-7	NA	0.02	<0.02
2G-1	NA	0.02	<0.02
2G-2	19	0.02	0.5
2G-3	19	0.02	2.0
2G-3 (Dup)	19	0.02	3.8
2G-4	17	0.02	7.7
2G-5	32	0.02	7.0
2G-6	NA	0.02	<0.02
2H-1	NA	0.02	<0.02
2H-2	36	0.02	7.9
2H-3	20	0.02	13
2H-4	25	0.02	9.9

TABLE 3-2 (continued)
 VERIFICATION SOIL SAMPLING RESULTS
 FOR AREA 2

September 3-4 and October 16, 1987			
Sample Description	Moisture Percent	PCBs ¹ , D.L. ²	mg/kg Concentration ³
2H-5	6.1	0.02	0.66
2H-6	NA	0.02	0.12
2H-6 (Dup)	NA	0.02	0.13
BL-1	NA	0.02	0.17
BL-2	0.15	0.02	0.5

Notes:

1. Only PCBs detected were Aroclor 1260.
2. D.L. is the detection limit.
3. Concentration of PCB detected by the laboratory analysis.
4. NA = Not analyzed.

TABLE 3-3

HUNTERS POINT ANNEX
TREASURE ISLAND NAVAL STATION

VERIFICATION SOIL SAMPLING RESULTS
FROM CENTRAL COAST ANALYTICAL SERVICES
FOR AREA 3

August 5-8, 1988			
Sample Description	Percent Moisture	PCBs ¹ D.L. ²	mg/kg Concentration ³
3A-9	7.8	1.0	<1.0
3A-11	7.4	1.0	<1.0
3A-13	22	1.0	<1.0
3B-6	4.4	1.0	<1.0
3B-8	13	1.0	4
3B10	22	1.0	9
3B10 (Dup)	NA	1.0	9
3B12	25	1.0	<1.0
3B16	8.0	1.0	<1.0
3B18	7.9	1.0	<1.0
3C-5	7.2	1.0	<1.0
3C-7	22	1.0	5
3C-9	20	1.0	<1.0
3C-11	20	1.0	<1.0
3C-13	22	1.0	6
3C-15	11	1.0	5
3C-17	6.7	1.0	<1.0
3C-19	30	1.0	<1.0
3D-4	6.5	1.0	<1.0
3D-6	19	1.0	2
3D-8	16	1.0	<1.0
3D-8 (Dup)	NA	1.0	<1.0
3D-10	14	1.0	3
3D-10 (Dup)	NA	1.0	3
3D-12	18	1.0	<1.0
3D-14	17	1.0	4
3D-16	12	1.0	<1.0
3D-18	11	1.0	<1.0

TABLE 3-3 (Continued)

HUNTERS POINT ANNEX
TREASURE ISLAND NAVAL STATION

VERIFICATION SOIL SAMPLING RESULTS
FROM CENTRAL COAST ANALYTICAL SERVICES
FOR AREA 3

August 5-8, 1988			
Sample Description	Percent Moisture	PCBs ¹ D.L. ²	mg/kg Concentration ³
3E-5	12	1.0	<1.0
3E-7	32	1.0	15
3E-7 (Dup)	NA	1.0	23
3E-9	21	1.0	14
3E-9 (Dup)	NA	1.0	17
3E-11	17	1.0	<1.0
3E-13	23	1.0	3
3E-15	13	1.0	<1.0
3E-17	13	1.0	<1.0
3F-4	8.0	1.0	<1.0
3F-6	20	1.0	7
3F-8	19	1.0	4
3F-10	14	1.0	2
3F-12	19	1.0	<1.0
3F-14	17	1.0	<1.0
3F-16	16	1.0	<1.0
3G-5	11	1.0	<1.0
3G-7	13	1.0	<1.0
3G-9	23	1.0	9
3G-11	18	1.0	2
3G-13	22	1.0	2
3G-15	5.3	1.0	<1.0
3H-6	7.1	1.0	<1.0
3H-8	17	1.0	<1.0
3H-10	15	1.0	<1.0
3H-12	9.0	1.0	<1.0
3H-14	12	1.0	1.0

TABLE 3-3 (Continued)

HUNTERS POINT ANNEX
TREASURE ISLAND NAVAL STATIONVERIFICATION SOIL SAMPLING RESULTS
FROM CENTRAL COAST ANALYTICAL SERVICES
FOR AREA 3

August 5-8, 1988			
Sample Description	Percent Moisture	PCBs ¹ D.L. ²	mg/kg Concentration ³
3J-9	11	1.0	<1.0
3J-11	17	1.0	2
3J-13	19	1.0	<1.0
3K-10	8.9	1.0	<1.0
3K-12	8.4	1.0	18
3K-12 (Dup)	NA	1.0	17

Notes:

1. Only PCBs detected were Aroclor 1260.
2. D.L. is the detection limit.
3. Concentration of PCBs detected in the laboratory analysis.
4. NA = Not analyzed.

APPENDIX B

PRELIMINARY SURVEY - SOIL SAMPLING RESULTS

Notes:

1. Boring and sample numbering correspond to circled numbers on report Figure 2-2
2. Table taken from reference 2
3. ANLAB was the certified laboratory used to analyze the soil samples

TABLE 1

HUNTER'S POINT PCB INVESTIGATION
OCTOBER 1986

BORING & SAMPLE NUMBER	DATE	TIME	McGRAW-EDISON FIELD KIT			ANLAB ANALYSIS		
			PROBE RESPONSE MV	ppm PCB IN SOIL		AROCLOR 1242	AROCLOR 1260	EPA 8010
				AROCLOR 1242	AROCLOR 1260			
1	15OCT86	940	156/167	<36	<16	X	X	X
2	15OCT86	955	124/117	44/60	20/26	X	26	X
3	15OCT86	1010	155	<36	<16	X	X	<0.05
4	15OCT86	1050	159	<36	<16	X	X	<0.05
5	15OCT86	1110	163	<36	<16	X	X	X
6	15OCT86	1640	156	<36	<16	X	<0.1	X
7	15OCT86	1100	159	<36	<16	X	X	X
10	15OCT86	1537	171	<36	<16	X	X	X
11	15OCT86	1500	169	<36	<16	X	X	X
12	15OCT86	1130	160	<36	<16	X	X	X
13	15OCT86	1430	163	<36	<16	X	X	X
14	15OCT86	1306	161	<36	<16	X	X	X
15	15OCT86	1233	165	<36	<16	X	X	X
16	16OCT86	1012	158	<36	<16	X	X	X
17	16OCT86	918	161	<36	<16	X	X	X

McGRAW-EDISON FIELD KIT

BORING & SAMPLE NUMBER	DATE	TIME	PROBE RESPONSE MV	ppm PCB IN SOIL		ANLAB ANALYSIS		
				AROCLOR 1242	AROCLOR 1260	AROCLOR 1242	AROCLOR 1260	EPA 8010
18	15OCT86	1546	171/170	<36	<16	X	X	X
19	15OCT86	1600	171	<36	<16	X	X	X
20	15OCT86	1314	149/141	<36	<16	X	X	X
21	15OCT86	1557	167	<36	<16	X	X	X
22	15OCT86	1155	157	<36	<16	X	X	X
23	16OCT86	1036	157	<36	<16	X	X	X
24	16OCT86	930	156/158	<36	<16	X	X	X
25	15OCT86	1145	165	<36	<16	X	X	X
26	16OCT86	1110	163	<36	<16	X	X	X
27	15OCT86	1200	163/160	<36	<16	X	X	X
28	16OCT86	1003	159	<36	<16	X	X	X
29	16OCT86	955	154	<36	<16	X	X	X
30	16OCT86	942	155	<36	<16	X	X	X
31	16OCT86	1031	158	<36	<16	X	X	X
32	16OCT86	1044	161	<36	<16	X	X	X
33	16OCT86	1051	158	<36	<16	X	X	X
35	16OCT86	1058	153	<36	<16	X	X	X
36	16OCT86	1105	155	<36	<16	X	X	X

McGraw-Edison Field Kit

Boring & Sample Number	Date	Time	Probe Response mV	ppm PCB in Soil		ANLAB ANALYSIS		
				AROCLOR 1242	AROCLOR 1260	AROCLOR 1242	AROCLOR 1260	EPA 8010
37	15OCT86	1035	156	<36	<16	X	X	X
38	15OCT86	1030	159	<36	<16	X	X	X
40	15OCT86	1225	156	<36	<16	X	X	X
41	16OCT86	1015	159	<36	<16	X	X	X
42	15OCT86	1210	158	<36	<16	X	X	X
43	15OCT86	1302	166	<36	<16	X	X	X
44	15OCT86	1255	160	<36	<16	X	X	X
45	16OCT86	1025	157/160	<36	<16	X	X	X
46	15OCT86	1246	168	<36	<16	X	X	X
2A	15OCT86	1615	-.019	>7260	>4200	X	810	X
2B	15OCT86	1620	166	<36	<16	X	X	X
2C	15OCT86	1631	168	<36	<16	X	X	X
2D(3ft)	16OCT86	813	158	<36	<16	X	X	X
2D(5ft)	16OCT86	815	162	<36	<16	X	X	X
2E(3ft)	16OCT86	854	168	<36	<16	X	X	X
2E(5ft)	16OCT86	856	159	<36	<16	X	X	X
2F(3ft)	16OCT86	843	165	<36	<16	X	X	X
2F(5ft)	16OCT86	845	158	<36	<16	X	X	X

McGRAW-EDISON FIELD KIT

BORING & SAMPLE NUMBER	DATE	TIME	PROBE RESPONSE MV	ppm PCB IN SOIL		ANLAB ANALYSIS		
				AROCLOR 1242	AROCLOR 1260	AROCLOR 1242	AROCLOR 1260	EPA 8010
2G(3ft)	16OCT86	832	134/136	<36	<16	X	7	X
2G(5ft)	16OCT86	835	156/155	<36	<16	X	2	X
2H(3ft)	16OCT86	823	164	<36	<16	X	X	X
2H(5ft)	16OCT86	825	159	<36	<16	X	X	X
2J(3ft)	16OCT86	900	163	<36	<16	X	X	X
2J(5ft)	16OCT86	903	163	<36	<16	X	X	X
2K	16OCT86	1208	163	<36	<16	X	X	X
2L	16OCT86	1200	157	<36	<16	X	X	X
2M	16OCT86	1315	160	<36	<16	X	X	X
2N	16OCT86	1219	162	<36	<16	X	X	X
2NO	16OCT86	1224	156/156	<36	<16	X	X	X
2P	16OCT86	1230	158	<36	<16	X	<0.1	X
2Q(6"-12")	16OCT86	1242	76	324	128	X	770	X
2Q(3ft)	16OCT86	1238	155	<36	<16	X	<0.1	X
2Q(5ft)	16OCT86	1239	----	<36	<16	X	<0.1	X
2R(3ft)	16OCT86	1300	23/34	2830/1800	1000/652	X	910	X
2R(5ft)	16OCT86	1302	35/37	1730/1594	628/580	X	180	X
2S	16OCT86	1247	142/143	<36	<16	X	<0.1	X

			McGRAW-EDISON FIELD KIT			ANLAB ANALYSIS		
BORING & SAMPLE NUMBER	DATE	TIME	PROBE	ppm PCB IN SOIL		AROCLOR 1242	AROCLOR 1260	EPA 8010
			RESPONSE MV	AROCLOR 1242	AROCLOR 1260			
2T	16OCT86	1259	153	<36	<16	X	X	X
2U	16OCT86	128	154	<36	<16	X	X	X

APPENDIX C

GROUND WATER STUDY
SOIL/GROUNDWATER DATA AND GEOLOGIC CROSS SECTION

APPENDIX C – GROUNDWATER STUDY
SOIL/GROUNDWATER DATA AND GEOLOGIC CROSS SECTION

THIS APPENDIX IS COMPLETE AS SUBMITTED.

FOR ADDITIONAL INFORMATION, CONTACT:

DIANE C. SILVA, RECORDS MANAGER
NAVAL FACILITIES ENGINEERING COMMAND, SOUTHWEST
1220 PACIFIC HIGHWAY
SAN DIEGO, CA 92132

TELEPHONE: (619) 532-3676
E-MAIL: diane.silva@navy.mil

TABLE 2-1

SUMMARY OF SUBSURFACE SOIL ANALYSES

<u>Well or Boring Number</u>	<u>Sample Date</u>	<u>Sample Depth</u>	<u>PCB Conc. (mg/kg)</u>
W1	1/7/87	10"-14"	28
W1	1/7/87	3'3"-3'9"	2.4
W1	1/7/87	5'0"-5'6"	32
W1	1/7/87	7'6"-8'0"	1
W1	1/7/87	10'0"-10'4"	4
W1	1/7/87	15'0"-15'6"	<0.1
W1'	1/7/87	16'0"-17'0"	<0.1
W1	1/7/87	20'0"-20'6"	24
Bl a	1/9/87	1'0" - 1'6"	1.2
Bl a	1/9/87	3'0" - 3'6"	2
W2	1/7/87	5'0" - 5'6"	5
W2	1/7/87	7'6" - 8'0"	4.3
W2	1/7/87	10'0"-10'6"	6
W2	1/7/87	11'0"-11'6"	<0.1
W2	1/7/87	12'6"-13'0"	0.6
W2	1/7/87	14'2"-14'8"	12
W3	1/8/87	2'6"- 3'0"	5.6
W3	1/8/87	0'6"- 15'0"	0.1
W4	1/8/87	2'6"- 3'0"	0.7
W4	1/8/87	0'6"- 14'8"	<0.1
W5	1/8/87	3'0"-3'6"	0.1
W5	1/8/87	0'6"-14'8"	<0.1
W6	1/9/87	3'0"-3'6"	<0.1
W6	1/9/87	0'6"-18'0"	0.1
W7	1/9/87	2'6"-2'10"	<0.1
W7	1/9/87	0.6"-15'0"	<0.1
B8	2/23/87	7'0"-7'6"	0.21
B9	2/23/87	2'6"-3'0"	0.05
B9	2/23/87	5'3"-5'9"	0.05
B9	2/23/87	3'6"-32'0"	0.05
B10	2/24/87	3'4"-3'10"	0.56
B10	2/24/87	5'2"-5'8"	0.06
B10	2/24/87	27'0"-27'6"	0.05

TABLE 2-5

SUMMARY OF GROUNDWATER AND SURFACE WATER ANALYSIS

<u>SAMPLE LOCATION</u>	<u>SAMPLING DATE</u>	<u>PCB CONC. (mg/l)</u>
W1	1/9/87	0.004
	3/1/87	<0.0005
W2	1/9/87	<0.001
	3/1/87	0.0007
W3	1/9/87	<0.001
	3/1/87	<0.0005
W4	1/9/87	<0.001
	3/1/87	<0.0005
W5	1/9/87	<0.001
	3/1/87	<0.0005
W6	1/12/87	<0.1
W7	1/12/87	<0.1
Catch Basin #1	1/9/87	0.004
Surface Seepage*	4/24/87	<0.0002

* Groundwater that filled excavation resulting from removal of transformer pad.

Elevation Above Mean Sea Level

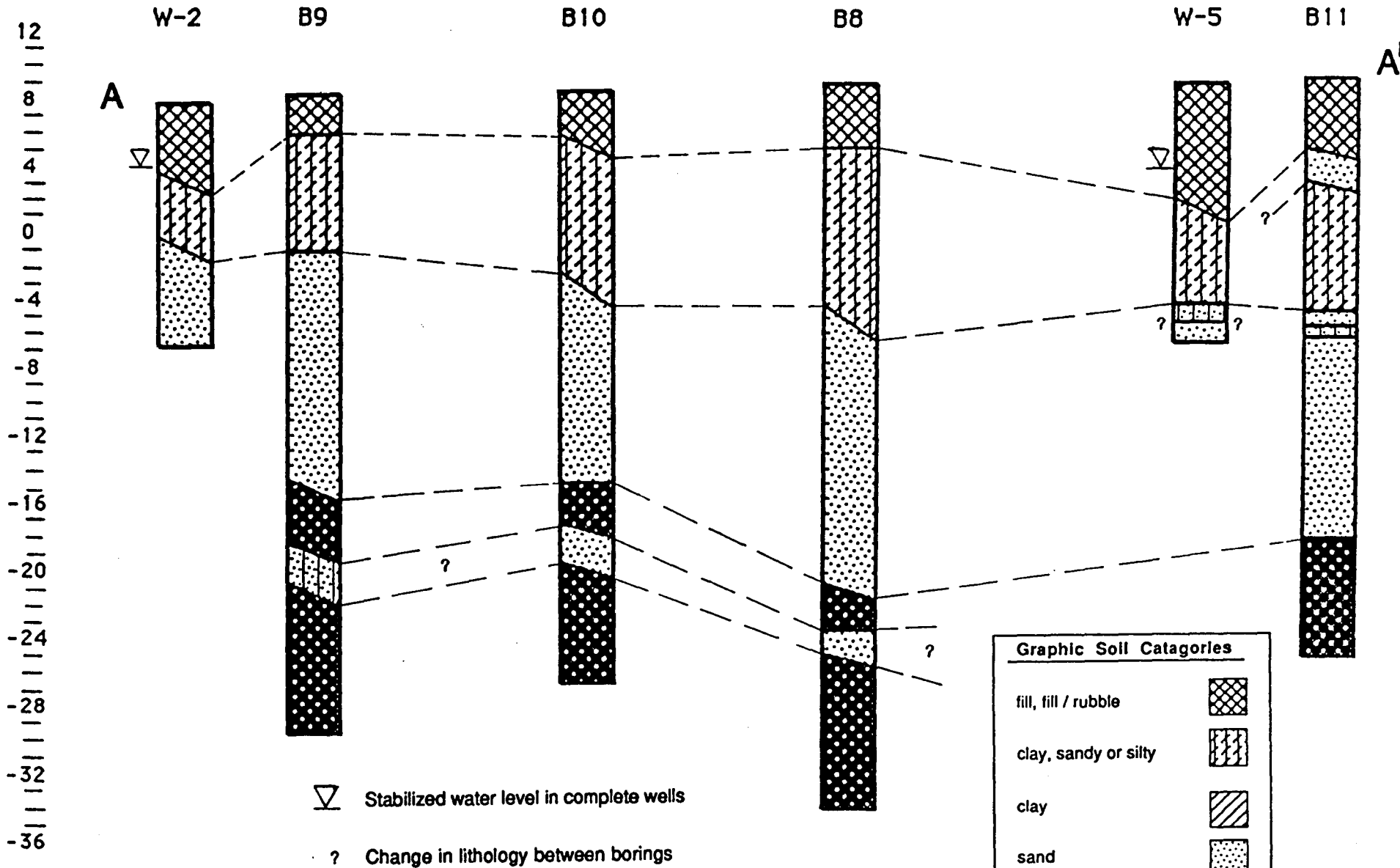


Figure 2-2a
Geologic Cross Section A - A'

Elevation Above Mean Sea Level

12
8
4
0
-4
-8
-12
-16
-20
-24
-28
-32
-36

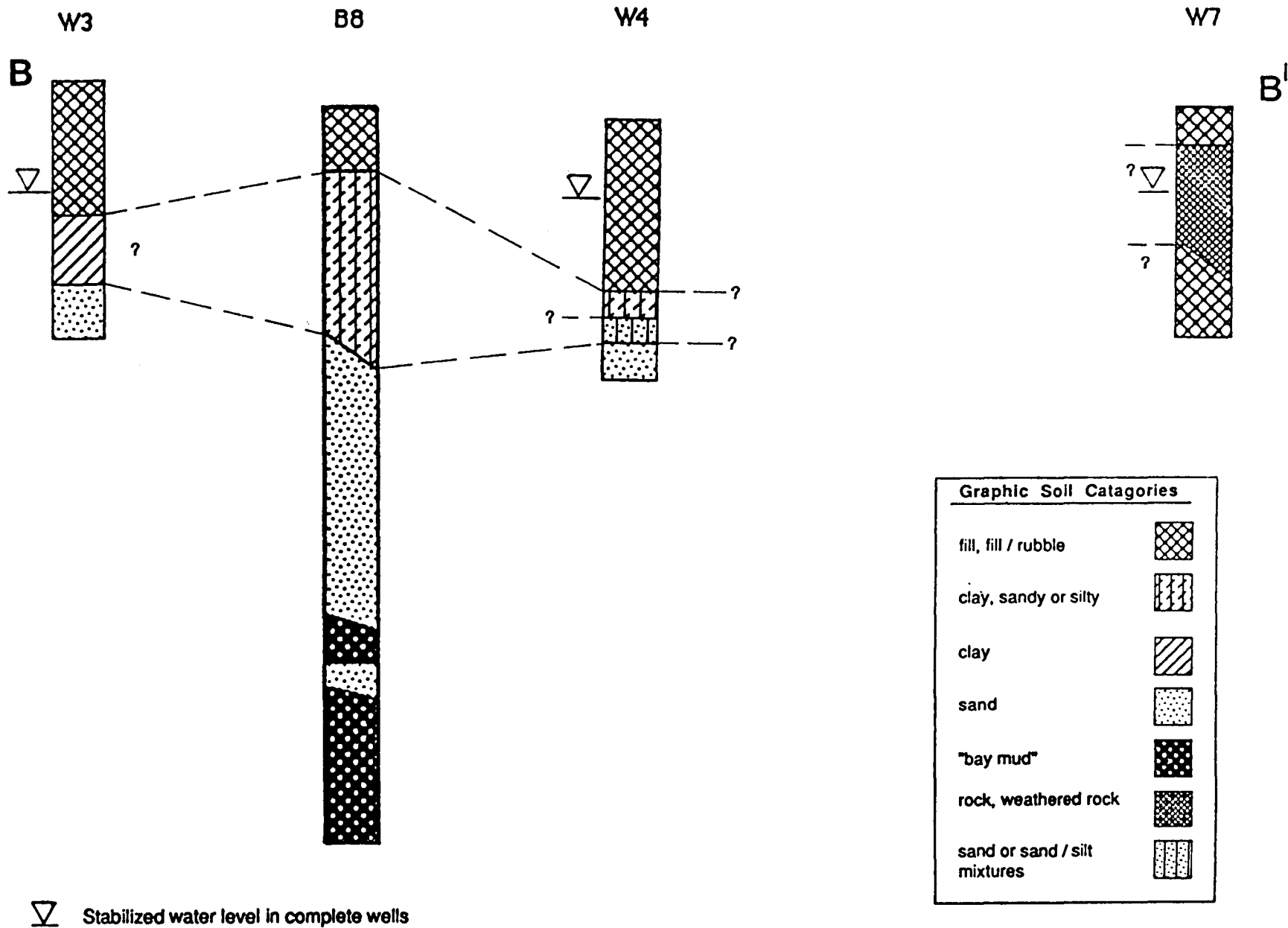


FIGURE 2-2b
Geologic Cross Section B - B'

APPENDIX D

VERIFICATION SOIL SAMPLING RESULTS
LABORATORY DATA REPORTS

D-1 AREA 1 (Interim Report)
D-2 AREA 2
D-3 AREA 3

· D-1 AREA 1 (Interim Report)

Central
Coast
Analytical
Services

Central Coast
Analytical Services
141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553

Lab Number: As Listed
Collected: 04/15/87
Received: 04/16/87 @ 1720
Tested: 04/19/87
Collected by: Dan Cutugno

ERM WEST
177 Botelho Drive
Suite 260
Walnut Creek, CA 94596

Sample Description:

US Navy Hunters Point, San Francisco,
CA, Soil Samples As Listed

REPORT

LAB NUMBER	SAMPLE DESCRIPTION	LEVEL FOUND - mg/kg
------------	--------------------	---------------------

POLYCHLORINATED BIPHENYLS
AROCOR 1260

EPA METHOD NUMBER-----
DETECTION LIMIT-----

608/8080
0.01

D-2680-2683	Composite A1, A2, B1, B2	71.
D-2684-2688	Composite C1, C2, D1, D2, D3	1.1
D-2689-2692	Composite E1, E2, E3, F1	2.1
D-2693-2696	Composite F2, F3, G1, G2	10.
D-2697-2700	Composite G4, H1, H2, H3	31.
QD-2697-2700	Duplicate of Above Sample	21.

Compositing was done as follows: Each soil sample (20g) was ground with a mortar and pestle then passed through a #10 sieve. The four ground and sieved discrete samples (20g each) were placed in a clean jar and mixed together. Approximately 20g of the resulting 80g composite was extracted using EPA Method 3550.

D2680COM.WR1/16
MH/ke

Respectfully submitted,
CENTRAL COAST ANALYTICAL SERVICES
Mary Havlicek
Mary Havlicek, Ph.D., President

Central
Coast
Analytical
Services

Central Coast
Analytical Services
141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553

Lab Number: As Listed
Collected: 04/15/87
Received: 04/16/87 @ 1720
Tested: 04/22/87
Collected by: Dan Cutugno

ERM WEST
1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596

Sample Description:

Project #400-30
US Navy Hunter's Point
Soil Samples As Listed

REPORT

LAB NUMBER	SAMPLE DESCRIPTION	LEVEL FOUND - mg/kg
------------	--------------------	---------------------

POLYCHLORINATED BIPHENYLS AROCOR 1260

EPA METHOD NUMBER-----	8080
------------------------	------

DETECTION LIMIT

000VD02A	Instrument Blank	0.02	<0.02
QH0421	Sonic Horn Extraction Blank	0.02	<0.02
D-2680	Soil A1	0.02	4.9
D-2681	Soil A2	0.2	46.
D-2682	Soil B1	0.02	32.
D-2683	Soil B2	0.2	190.
D-2697	Soil G4	0.2	35.
D-2699	Soil H3	0.2	3.6
QD-2699	Duplicate of Above	0.2	6.6
D-2700	Soil H1	0.2	2.0

D2680ERM.WR1/17
MH/ke

Respectfully submitted,
CENTRAL COAST ANALYTICAL SERVICES
Mary Havlicek
Mary Havlicek, Ph.D., President

Central
Coast
Analytical
Services

Central Coast
Analytical Services
141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553

Lab Number: As Listed
Collected: 04/15/87
Received: 04/16/87
Tested: 05/01/87
Collected by: Dan Cutugno

ERM WEST
1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596

Sample Description:

US Navy, Hunters Point
Soil Samples As Listed

REPORT

LAB NUMBER	SAMPLE DESCRIPTION	LEVEL FOUND -mg/kg
------------	--------------------	--------------------

POLYCHLORINATED BIPHENYLS
AROCOR 1260

EPA METHOD NUMBER-----
DETECTION LIMIT-----

608/8080
0.1

D-2693	Soil F-2	3.5
D-2694	Soil F-3	1.0
D-2695	Soil G-1	4.2
D-2696	Soil G-2	0.4

D2693ERM.WR1/21
MH/ke

Respectfully submitted,
CENTRAL COAST ANALYTICAL SERVICES
Mary Havlicek
Mary Havlicek, Ph.D., President

Central
Coast
Analytical
Services

Central Coast
Analytical Services
141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553

Lab Number: As Listed
Collected: 05/18/87
Received: 05/19/87 @ 0900
Tested: 05/21/87
Collected by: Dennis Miller

ERM-WEST
1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596

Sample Description:

Job #40030 US Navy, Hunter's Point
Soil Samples As Listed

REPORT

LAB NUMBER	SAMPLE DESCRIPTION	LEVEL FOUND - mg/kg
------------	--------------------	---------------------

POLYCHLORINATED BIPHENYLS

AROCLOR 1260

608/8080

0.01

EPA METHOD NUMBER-----
DETECTION LIMIT-----

D-3600	X1 @ 2'6"	0.30
D-3601	X2 @ 2'2"	0.49
D-3602	X3 @ 2'4"	0.03
D-3603	X4 @ 2'0"	0.08
D-3604	X5 @ 2'0"	0.02
D-3605	X6 @ 2'2"	0.02
D-3606	X7 @ 1'11"	0.04
D-3607	X8 @ 1'10"	<0.01
D-3608	X9 @ 2'1"	0.04
D-3609	X10 @ 1'9"	0.10
D-3610	X11 @ 2'0"	0.04
D-3611	X12 @ 2'4"	0.05
QD-3611	DUPLICATE	0.05
D-3612	X13 @ 2'0"	0.25
D-3613	X14 @ 2'0"	0.10
D-3614	X15 @ 2'0"	0.08
D-3615	X16 @ 2'6"	0.16

D3600PCB.WR1
MH/ke

Respectfully submitted,
CENTRAL COAST ANALYTICAL SERVICES
Mary Havlicek
Mary Havlicek, Ph.D., President

D-2 AREA 2

Central
Coast
Analytical
Services

Central Coast
Analytical Services
141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553

Lab Number: As Listed
Collected: 09/04/87
Received: 09/05/87
Tested: As Listed
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Drive
Suite 260
Walnut Creek, CA 94523

Sample Description:

Hunter's Point, U.S. Navy
Project #400-36
Samples As Listed

REPORT

LAB NUMBER	SAMPLE DESCRIPTION	PERCENT MOISTURE	LEVEL FOUND -
EPA METHOD NUMBER-----		Percent	POLYCHLORINATED
DETECTION LIMIT-----		160.3	BIPHENYLS
DATE TESTED/ANALYST-----		0.005	AROCLOR 1260
		9/11/87/LD	mg/kg (Wet Weight)
			8080
			0.02
			9/08-15/87/ARP
D-6950	2A-2	3.2	5.0
D-6951	2A-3	7.1	2.2
D-6952	2A-4	2.7	0.61
D-6953	2B-2	16.	16.
D-6954	2B-3	7.2	0.44
OD-6954	DUPLICATE	7.1	---
D-6955	2B-4	5.1	1.0
D-6956	2B-5	3.8	0.60
D-6957	2C-3	12.	3.2

D6950ERM.WR1
MH/ko

Respectfully submitted,
CENTRAL COAST ANALYTICAL SERVICES
Mary Havlicek
Mary Havlicek, Ph.D., President

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141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553

Lab Number: As Listed
Collected: 09/03/87
Received: 09/04/87
Tested: 09/07/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Drive
Suite 260
Walnut Creek, CA 94598

Sample Description:
U.S. Navy Hunter's Point
Project #400-36

REPORT

LAB NUMBER	SAMPLE DESCRIPTION	TOTAL LEVEL FOUND- milligrams/kg (Wet Weight)
POLYCHLORINATED BIPHENYLS		
EPA METHOD NUMBER-----		8080
DETECTION LIMIT-----		0.02
DATE TESTED/ANALYST-----		9/7/87/ARP
D-6886	2H-1 @ 1'6"-2'0"	<0.02
D-6887	2G-1 @ 1'8"-2'3"	<0.02
D-6888	2F-1 @ 1'5"-2'0"	<0.02
D-6889	2D-5 @ 2'3"-2'8"	<0.02
D-6890	2B-6 @ 1'6"-2'0"	<0.02
D-6891	2C-7 @ 2'7"-3'1"	<0.02
D-6892	2D-6 @ 1'6"-2'0"	<0.02
D-6893	2E-7 @ 1'8"-2'2"	<0.02
D-6894	2F-7 @ 1'10"-2'4"	<0.02
D-6895	2G-6 @ 2'4"-2'10"	<0.02
D-6896	2H-6 @ 1'4"-1'10"	0.12
QJ-6896	DUPLICATE	0.13
D-6897	2B-1 @ 1'8"-2'2"	<0.02
D-6898	2C-1 @ 1'6"-2'0"	0.04
D-6899	BL-1 @ 1'6"-2'0"	0.17

D6886ERM.WR1
MH/ke

Respectfully submitted,
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Mary Havlicek, Ph.D., President

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San Luis Obispo, California 93401
(805) 543-2553

Lab Number: As Listed
Collected: 09/04/87
Received: 09/05/87
Tested: As Listed
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Drive
Suite 260
Walnut Creek, CA 94523

Sample Description:

Hunter's Point, U.S. Navy
Project #400-36
Samples As Listed

REPORT

LAB NUMBER

SAMPLE DESCRIPTION

LEVEL FOUND -

PERCENT MOISTURE

POLYCHLORINATED
BIPHENYLS

AROCLOR 1260

Percent

mg/kg (Wet Weight)

EPA METHOD NUMBER-----

160.3

8080

DETECTION LIMIT-----

0.005

0.02

DATE TESTED/ANALYST-----

9/11/87/LD

9/08-15/87/ARP

D-6958	2C-4	25.	15.
D-6959	2D-1	30.	3.5
D-6960	2D-2	41.	21.
D-6961	2D-3	31.	4.3
D-6962	2D-4	21.	2.2
D-6963	2D-5	1.2	2.2
D-6964	2E-2	41.	36.
D-6965	2E-3	25.	59.
D-6966	2E-5	20.	4.6
QD-6966	DUPLICATE	20.	4.2
D-6967	2E-6	17.	0.60
D-6968	2F-2	25.	3.4
D-6969	2F-3	30.	9.6

D6958ERM.WR1
MH/ke

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(805) 543-2553

Lab Number: As Listed
Collected: 9/04/87
Received: 9/05/87
Tested: As Listed
Collected by: Richard Knapp

ERM-WEST
1777 Botelho Drive
Suite 260
Walnut Creek, CA 94523

Sample Description:

Hunter's Point, U.S. Navy
Project #400-36
Samples As Listed

REPORT

LAB NUMBER	SAMPLE DESCRIPTION	PERCENT MOISTURE	LEVEL FOUND -
			POLYCHLORINATED BIPHENYLS AROCOR 1260 mg/kg (Wet Weight)
EPA METHOD NUMBER-----		Percent	
DETECTION LIMIT-----		16.3	8080
DATE TESTED/ANALYST-----		0.005	0.02
		9/11/87/LD	9/11/87/ARP
D-6970	2F-4	17.	8.4
D-6971	2F-5	23.	0.73
D-6972	2F-6	1.6	0.6
D-6973	2G-2	19.	0.50
D-6974	2G-3	19.	2.0
00-6974	DUPLICATE	19.	3.8
D-6975	2G-5	32.	7.0
D-6976	2H-2	36.	7.9
D-6977	2H-3	20.	13.0
D-6978	2H-4	25.	9.9
D-6979	2H-5	6.1	0.66
D-6980	BL-2	0.15	0.50

D6970ERM.WR1
MH/ke

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Mary Havicek, Ph.D., President

c[<1[10m AIR, WATER and HAZARDOUS WASTE LABORATORY CERTIFIED by CALIFORNIA DEPT of PUBLIC HE

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Lab Number: As Listed
Collected: 10/16/87
Received: 10/17/87
Tested: As Listed
Collected by: D. Catugno

ERM-WEST
1777 Botelho Drive
Suite 260
Walnut Creek, CA 94598

Sample Description:

US Navy, Hunter's Point
Project #400-36, S.F.

REPORT

LAB NUMBER

SAMPLE DESCRIPTION

LEVEL FOUND-

EPA METHOD NUMBER-----
DETECTION LIMIT-----
DATE TESTED/ANALYST-----

PERCENT MOISTURE %	PERCENT SOLID %
160.3	160.3
0.005	0.005
10/21/87/RDM	10/21/87/RDM

D-8233	Sample #2C-5A	15.	85.
D-8234	Sample #2C-5B	21.	79.
D-8235	Sample #2C-5C	19.	81.
D-8236	Sample #2E-4	19.	81.
D-8237	Sample #2D-2	24.	76.
D-8238	Sample #2E-3B	18.	84.
D-8239	Sample #2E-3A	20.	80.
D-8240	Sample #2E-2A	5.	95.
D-8241	Sample #2E-2	18.	82.

D0233ERM.WR1
MH/ke

Respectfully submitted,
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EPA METHOD 608/8080 - PCB'S

Lab Number: D-7239
Collected: 9/04/78
Received: 9/17/87
Tested: 9/21/87
Collected by: Dan Catugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
U.S Navy Hunters Point
HML#500, 2A-1, Xn-6.8; Xs-20
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.2	not found
PCB 1221	0.2	not found
PCB 1232	0.2	not found
PCB 1242	0.2	not found
PCB 1248	0.2	not found
PCB 1254	0.2	not found
PCB 1260	0.2	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 9/18/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

D7239PC.WR1
MH/TK/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: D-7245
Collected: 9/04/78
Received: 9/17/87
Tested: 9/21/87
Collected by: Dan Catugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
U.S Navy Hunters Point
HML#512,2C-2, %m=30;%s=38
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.02	not found
PCB 1221	0.02	not found
PCB 1232	0.02	not found
PCB 1242	0.02	not found
PCB 1248	0.02	not found
PCB 1254	0.02	not found
PCB 1260	0.02	25.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 9/18/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

D7245PC.WR1
MH/TK/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lob Number: D-7242
Collected: 9/04/78
Received: 9/17/87
Tested: 9/21/87
Collected by: Dan Catugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
U.S Navy Hunters Point
HML#509,2C-5, %m<0.1;%s=17
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.02	not found
PCB 1221	0.02	not found
PCB 1232	0.02	not found
PCB 1242	0.02	not found
PCB 1248	0.02	not found
PCB 1254	0.02	not found
PCB 1260	0.02	51.

Compounds listed as "not found", would have been reported if present
at or above the listed detection limits. Sample was extracted 9/18/87

Respectfully submitted,

Mary Havlicek

Mary Havlicek, Ph.D.

D7242PC.WR1
MH/GH/ARP/ARP

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EPA METHOD 808/8080 - PCB'S

Lab Number: D-8552
Collected: 10/16/87
Received: 10/26/87
Tested: 11/05/87
Collected by: Dan Cutugno

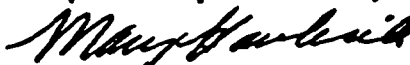
ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
2c-5, Hunter's Point, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.1	not found
PCB 1221	0.1	not found
PCB 1232	0.1	not found
PCB 1242	0.1	not found
PCB 1248	0.1	not found
PCB 1254	0.1	not found
PCB 1260	0.1	1.3

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/28/87

Respectfully submitted,



Mary Havlicek, Ph.D.

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EPA METHOD 808/8080 - PCB'S

Lab Number: QT-8241
Collected: 10/18/87
Received: 10/17/87
Tested: 10/22/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2C-5A
PROJECT #400-38

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	2.7

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

QT8241pc.wr1
PH/JK/ARP/ARP

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EPA METHOD 606/8060 - PCB'S

Lab Number: D-8233
Collected: 10/16/87
Received: 10/16/87
Tested: 10/22/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2C-5A
PROJECT #400-30

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	2.4

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87.

Respectfully submitted,

Mary Havlicek
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EPA METHOD 808/8080 - PCB's

Lab Number: QQ-8241
Collected: 10/16/87
Received: 10/17/87
Tested: 10/23/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2C-5A
PROJECT #400-38

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	2.9

Compounds listed as "not found" would have been reported if present at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek

Mary Havlicek, Ph.D.

OCT 26 '87 15:14 C C R S

P04

c[1[15m AIR, WATER and HAZARDOUS WASTE LABORATORY CERTIFIED by CALIFORNIA DEPT of HEALTH SERVICE

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EPA METHOD 808/8080 - PCB'S

Lab Number: D-8234
Collected: 10/16/87
Received: 10/17/87
Tested: 10/22/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2C-58
PROJECT #458-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	11.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87.

Respectfully submitted,

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D8234PC.WR1
MH/JK/ARP/ARP

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Lab Number: D-8235
Collected: 10/16/87
Received: 10/16/87
Tested: 10/22/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2C-5C
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	1.5

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87 .

Respectfully submitted,

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Lab Number: D-7241
Collected: 9/04/78
Received: 9/17/87
Tested: 9/21/87
Collected by: Don Catugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
U.S Navy Hunters Point
HML#502,2C-6, %m=4.7;%s=19
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.02	not found
PCB 1221	0.02	not found
PCB 1232	0.02	not found
PCB 1242	0.02	not found
PCB 1248	0.02	not found
PCB 1254	0.02	not found
PCB 1260	0.02	0.65

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 9/18/87

Respectfully submitted,



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D7241PC.WR1
MH/TK/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: D-8237
Collected: 10/16/87
Received: 10/17/87
Tested: 10/22/87
Collected by: Don Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
20-2
PROJECT #400-30

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	16.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87.

Respectfully submitted,

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Mary Havlicek, Ph.D.

D8237PC.WR1
MH/JK/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: QD-7240
Collected: 9/04/78
Received: 9/17/87
Tested: 9/21/87
Collected by: Dan Catugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
U.S Navy Hunters Point
HML#502,2E-1, Km-5.8; %s=15
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.2	not found
PCB 1221	0.2	not found
PCB 1232	0.2	not found
PCB 1242	0.2	not found
PCB 1248	0.2	not found
PCB 1254	0.2	not found
PCB 1260	0.2	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 9/18/87

Respectfully submitted,

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QD7240PC.WR1
MH/TK/ARP/ARP

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Lab Number: D-7240
Collected: 9/04/78
Received: 9/17/87
Tested: 9/21/87
Collected by: Dan Catugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
U.S. Navy Hunters Point
HML#502, 2E-1, %m=5.8; %s=15
PROJECT #400-36

s/be HML#501

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.2	not found
PCB 1221	0.2	not found
PCB 1232	0.2	not found
PCB 1242	0.2	not found
PCB 1248	0.2	not found
PCB 1254	0.2	not found
PCB 1260	0.2	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 9/18/87

Respectfully submitted,

Mary Havlicek
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D7240PC.WR1
MH/TK/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: QO-8241
Collected: 10/16/87
Received: 10/16/87
Tested: 10/21/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2E-2
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	3.0

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

QO8241PC.WR1
MH/GH/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: D-8241
Collected: 10/16/87
Received: 10/18/87
Tested: 10/21/87
Collected by: Don Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2E-2
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	3.0

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

OCT 26 '87 15:18 C C R S

P11

c[<1[10m AIR, WATER and HAZARDOUS WASTE LABORATORY CERTIFIED by CALIFORNIA DEPT of HEALTH SE

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EPA METHOD 608/8080 - PCB'S

Lab Number: D-8240
Collected: 10/16/87
Received: 10/16/87
Tested: 10/21/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2E-2A
PROJECT #400-30

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	2.2

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

D8240PC.WR1
MH/JJK/ARP/ARP

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EPA METHOD 808/8080 - PCB'S

Lab Number: D-8553
Collected: 10/18/87
Received: 10/28/87
Tested: 11/05/87
Collected by: D. Cutugno

ERM-WEST
1777 Botelho Dr., Suite 280
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
2E-3, Hunter's Point, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	7.9

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/28/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

D8553pc.wr1/3
MH/gh/arp/sc

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EPA METHOD 600/8080 - PCB'S

Lab Number: D-8239
Collected: 10/16/87
Received: 10/16/87
Tested: 10/21/87
Collected by: Don Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2E-3A
PROJECT #400-38

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	15.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

D8239PC.WR1
MH/GH/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: D-8238
Collected: 10/16/87
Received: 10/16/87
Tested: 10/21/87
Collected by: Don Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2E-38
PROJECT #400-38

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	0.79

Compounds listed as "not found" would have been reported if present at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

OCT 26 '87 15:15 C C R S

P06

c[<1[15m AIR, WATER and HAZARDOUS WASTE LABORATORY CERTIFIED by CALIFORNIA DEPT of HEALTH SERV

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EPA METHOD 808/8080 - PCB'S

Lab Number: D-8236
Collected: 10/16/87
Received: 10/16/87
Tested: 10/22/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point
2E-4
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	8.9

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87.

Respectfully submitted,

Mary Havlicek

Mary Havlicek, Ph.D.

D8236PC.WR1
PHI/JK/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: D-7243
Collected: 9/04/78
Received: 9/17/87
Tested: 9/21/87
Collected by: Dan Catugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
U.S Navy Hunters Point
HML#510, 2E-4, Xm-22; Xs-40
PROJECT #400-38

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.02	not found
PCB 1221	0.02	not found
PCB 1232	0.02	not found
PCB 1242	0.02	not found
PCB 1248	0.02	not found
PCB 1254	0.02	not found
PCB 1260	0.02	24.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 9/18/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

D7243PC.WR1
MH/TK/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: QD-7243
Collected: 9/04/78
Received: 9/17/87
Tested: 9/21/87
Collected by: Dan Catugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
U.S Navy Hunters Point
HML#510, 2E-4, Sm-22; Ss-40
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.02	not found
PCB 1221	0.02	not found
PCB 1232	0.02	not found
PCB 1242	0.02	not found
PCB 1248	0.02	not found
PCB 1254	0.02	not found
PCB 1260	0.02	25.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 9/18/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

QD-7243PC.WR1
MH/TK/ARP/ARP

OCT 26 '87 15:16 U C H S

P07

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Lab Number: QS-8236
Collected: 10/16/87
Received: 10/16/87
Tested: 10/23/87
Collected by: Dan Cutugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94598

SAMPLE DESCRIPTION:
Erm West; Navy, Hunter's Point, 2E-4,
Spiked at 4.4 ppm
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	14.0

*Recovery of spike = 118%

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

QS8236PC.WR1
MH/GH/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: D-7244
Collected: 9/04/78
Received: 9/17/87
Tested: 9/21/87
Collected by: Dan Catugno

ERM-WEST
1777 Botelho Dr., Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
U.S Navy Hunters Point
HML#511,2G-4, Sm=17;Ss=48
PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.02	not found
PCB 1221	0.02	not found
PCB 1232	0.02	not found
PCB 1242	0.02	not found
PCB 1248	0.02	not found
PCB 1254	0.02	not found
PCB 1260	0.02	7.7

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 9/18/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

D7244PC.WR1
MH/TK/ARP/ARP

OCT 26 '87 15:26 C C R S

P21

- c[<1[10m AIR, WATER and HAZARDOUS WASTE LABORATORY CERTIFIED by CALIFORNIA DEPT of HEALTH SERVICE

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EPA METHOD 608/8080 - PCB'S

Lab Number: QL-1002
Collected:
Received:
Tested: 10/23/87
Collected by:

CCAS

SAMPLE DESCRIPTION:
Liq;Liq Extraction Blank

Compound Analyzed	Detection Limit micrograms/Kg	Concentration micrograms/Kg
PCB 1016	0.2	not found
PCB 1221	0.2	not found
PCB 1232	0.2	not found
PCB 1242	0.2	not found
PCB 1248	0.2	not found
PCB 1254	0.2	not found
PCB 1260	0.2	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

QL1002PC.WR1
MH/GH/ARP/ARP

c[<1[10m AIR, WATER and HAZARDOUS WASTE LABORATORY CERTIFIED by CALIFORNIA DEPT of HEALTH SJ

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Lab Number: B-10237
Collected:
Received:
Tested: 10/23/87
Collected by:

CCAS

SAMPLE DESCRIPTION:
Instrument Blank

Compound Analyzed	Detection Limit micrograms/Kg	Concentration micrograms/Kg
PCB 1016	0.2	not found
PCB 1221	0.2	not found
PCB 1232	0.2	not found
PCB 1242	0.2	not found
PCB 1248	0.2	not found
PCB 1254	0.2	not found
PCB 1260	0.2	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

B1023PC.WR1
MH/GH/ARP/ARP

c[<1[10m AIR, WATER and HAZARDOUS WASTE LABORATORY CERTIFIED by CALIFORNIA DEPT of HEALTH SERV

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EPA METHOD 608/8080 - PCB'S

Lab Number: QL-1555
Collected: 10/05/87
Received: 10/05/87
Tested: 10/22/87
Collected by: na

CCAS

SAMPLE DESCRIPTION:
Liquid-Liquid Extraction Blank

Compound Analyzed	Detection Limit micrograms/L	Concentration micrograms/L
PCB 1016	0.2	not found
PCB 1221	0.2	not found
PCB 1232	0.2	not found
PCB 1242	0.2	not found
PCB 1248	0.2	not found
PCB 1254	0.2	not found
PCB 1260	0.2	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/05/87.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

QL1005pc.wr1
MH/GH/ARP/ARP

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P18

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EPA METHOD 808/8080 - PCB'S

Lab Number: B-10227
Collected: no
Received: no
Tested: 10/22/87
Collected by: no

CCAS

SAMPLE DESCRIPTION:
Instrument Blank

Compound Analyzed	Detection Limit micrograms/L	Concentration micrograms/L
PCB 1016	0.2	not found
PCB 1221	0.2	not found
PCB 1232	0.2	not found
PCB 1242	0.2	not found
PCB 1248	0.2	not found
PCB 1254	0.2	not found
PCB 1260	0.2	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

B10227pc.wr1
MH/JK/ARP/ARP

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EPA METHOD 600/8-80-01 - PCB'S

Lab Number: B10217
Collected:
Received:
Tested: 10/21/87
Collected by:

CCAS

SAMPLE DESCRIPTION:
Instrument Blank

PROJECT #400-38

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

B10217PC.WR1
MH/JJK/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: B-89217
Collected:
Received:
Tested: 9/21/87
Collected by:

CCAS

SAMPLE DESCRIPTION:
Instrument Blank

PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.02	not found
PCB 1221	0.02	not found
PCB 1232	0.02	not found
PCB 1242	0.02	not found
PCB 1248	0.02	not found
PCB 1254	0.02	not found
PCB 1260	0.02	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

B89217PC.WR1
MH/GH/ARP/ARP

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EPA METHOD 608/8080 - PCB'S

Lab Number: QH10207
Collected:
Received:
Tested: 10/21/87
Collected by:

CCAS

SAMPLE DESCRIPTION:
Sonic Horn Extraction Blank

PROJECT #400-36

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.05	not found
PCB 1221	0.05	not found
PCB 1232	0.05	not found
PCB 1242	0.05	not found
PCB 1248	0.05	not found
PCB 1254	0.05	not found
PCB 1260	0.05	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 10/20/87

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.

QH10207PC.WR1
MH/JJK/ARP/ARP

D-3 AREA 3

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7681
Collected: 08/08/88 @ 1141
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

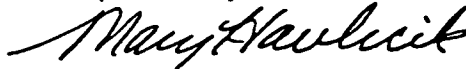
SAMPLE DESCRIPTION:
3A-9 (1.5 to 2.0 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 7.8%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7681pc.wr1/35
MH/bl/jl/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7679
Collected: 08/08/88 @ 1218
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3A-11 (2.0 to 2.5 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 7.4%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7679pc.wr1/36
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7678
Collected: 08/08/88 @ 1205
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

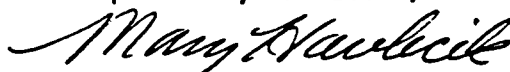
SAMPLE DESCRIPTION:
3A-13 (2.0 to 2.5 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 22%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7678pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7680
Collected: 08/08/88 @ 1228
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
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Walnut Creek, CA 94596

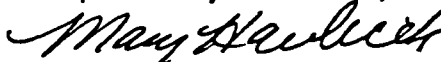
SAMPLE DESCRIPTION:
3B-6 (2.0 to 2.5 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 4.4%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7680pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7572
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

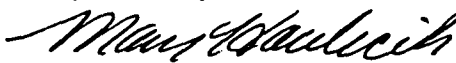
SAMPLE DESCRIPTION:
3B-8, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	4.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 13%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7572pc.wr1/35
MH/bl/sc/jl

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141 Suburban Road , Suite C-4
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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7573
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

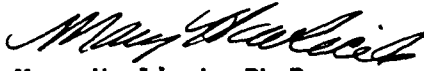
SAMPLE DESCRIPTION:
3B-10, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	9.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 22%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

MSD #5
E7573pc.wr1/35
MH/bl/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7573
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09 & 24/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

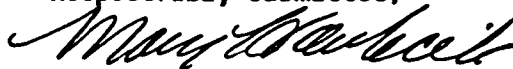
SAMPLE DESCRIPTION:
3B-10, Soil
Duplicate Analysis of Same Extract

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	9.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 22%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
08/26/88
E7573pc2.wr1/37
MH/als/sc/sc

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(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7574
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

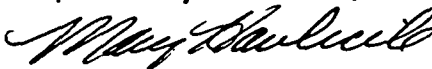
SAMPLE DESCRIPTION:
3B-12, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 13%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7574pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7677
Collected: 08/08/88 @ 1153
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3B-14 (2.0 to 2.5 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/09/88.

Percent Moisture: 25%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek

Mary Havlicek, Ph.D.
President

MSD #5
E7677pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7676
Collected: 08/08/88 @ 1142
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596


SAMPLE DESCRIPTION:
3B-16 (2.0 to 2.5 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/09/88.

Percent Moisture: 8.0%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

MSD #5
E7676pc.wr1/35
MH/bl/jl/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7675
Collected: 08/08/88 @ 1129
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3B-18 (2.0 to 2.5 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/09/88.

Percent Moisture: 7.9%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
E7675pc.wr1/35
MH/bl/jl/sc

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(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7682
Collected: 08/08/88 @ 1342
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

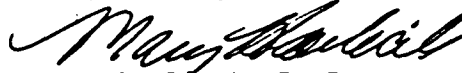
SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3C-5 (1.5 to 2.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 7.2%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

ECD
E7682pc.wr1/36
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7568
Collected: 08/05/88
Received: 08/07/88
Tested: 08/11/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

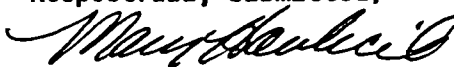
SAMPLE DESCRIPTION:
3C-7, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	5.

Compounds listed as "not found" would have been reported if present at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 22%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7568pc.wr1/35
MH/bl/sc/sc

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Lab Number: E-7570
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

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1777 Botelho Dr.
Walnut Creek, CA 94596

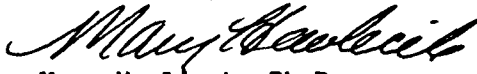
SAMPLE DESCRIPTION:
3C-9, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 20%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

MSD #5
E7570pc.wr1/35
MH/bl/sc/jl

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(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7571
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3C-11, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 20%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
E7571pc.wr1/35
MH/bl/sc/jl

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(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7575
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

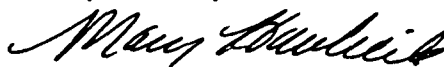
SAMPLE DESCRIPTION:
3C-13, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	6.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 22%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7575pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7576
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3C-15, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	5.

Compounds listed as "not found" would have been reported if present at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 11%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7576pc.wr1/35
MH/bl/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7674
Collected: 08/08/88 @ 1113
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

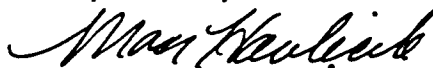
SAMPLE DESCRIPTION:
3C-17 (2.0 to 2.5 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/09/88.

Percent Moisture: 6.7%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7674pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7673
Collected: 08/08/88 @ 1056
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3C-19 (2.0 to 2.5 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/09/88.

Percent Moisture: 30%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
E7673pc.wr1/35
MH/bl/jl/sc

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Lab Number: E-7683
Collected: 08/08/88 @ 1352
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

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Suite 260
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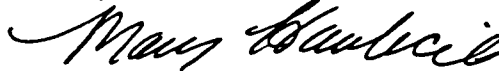
SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3D-4 (2.5 to 3.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 6.5%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

ECD
E7683pc.wr1/36
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7567
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

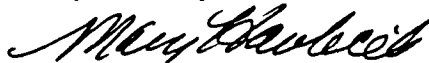
SAMPLE DESCRIPTION:
3D-6, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	2.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 19%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7567pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7569
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

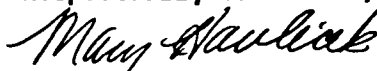
SAMPLE DESCRIPTION:
3D-8, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 16%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7569pc.wr1/35
MH/bl/sc/jl

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7569
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09 & 24/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
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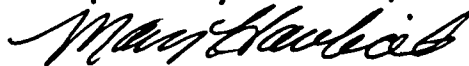
SAMPLE DESCRIPTION:
3D-8, Soil
Duplicate Analysis of Same Extract

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 16%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
08/26/88
E7569pc2.wr1/37
MH/als/sc/j1

Central
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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7566
Collected: 08/05/88
Received: 08/07/88
Tested: 08/11/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3D-10, Soil

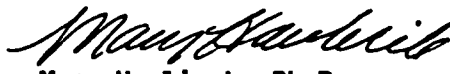
REVISED REPORT

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	3.

Compounds listed as "not found" would have been reported if present at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 14%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

MSD #5
E7566pc.wr1/35
MH/bl/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7566
Collected: 08/05/88
Received: 08/07/88
Tested: 08/11 & 24/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3D-10, Soil
Duplicate Analysis of Same Extract

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	3.

Compounds listed as "not found" would have been reported if present at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 14%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

MSD #5/ECD
08/26/88
E7566pc2.wr1/37
MH/als/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7565
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

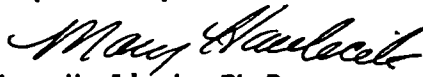
SAMPLE DESCRIPTION:
3D-12, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 18%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

MSD #5
E7565pc.wr1/35
MH/bl/sc/sc

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San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7577
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

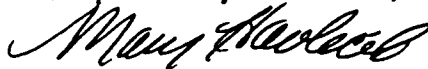
SAMPLE DESCRIPTION:
3D-14, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	4.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 17%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7577pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7699
Collected: 08/08/88 @ 1715
Received: 08/09/88 @ 1800
Tested: 08/12/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3D-16 (1.5 to 2.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 12%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
E7699pc.wr1/36
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7672
Collected: 08/08/88 @ 1038
Received: 08/09/88
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3D-18 (1.5 to 2.0 Feet), Soil Grab

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/09/88.

Percent Moisture: 11%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
E7672pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7684
Collected: 08/08/88 @ 1406
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

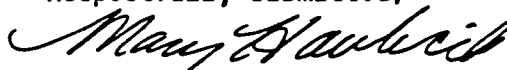
SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3E-5 (2.0 to 2.5 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 12%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

ECD
E7684pc.wr1/36
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7564
Collected: 08/05/88
Received: 08/07/88
Tested: 08/11/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

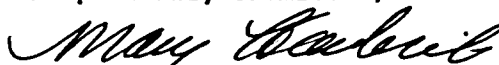
SAMPLE DESCRIPTION:
3E-7, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	15.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 32%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7564pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7564
Collected: 08/05/88
Received: 08/07/88
Tested: 08/11 & 24/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

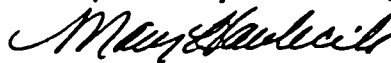
SAMPLE DESCRIPTION:
3E-7, Soil
Duplicate Analysis of Same Extract

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	23.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 32%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
08/26/88
E7564pc2.wr1/37
MH/als/sc/sc

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San Luis Obispo, California 93401
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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7563
Collected: 08/05/88
Received: 08/07/88
Tested: 08/11/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

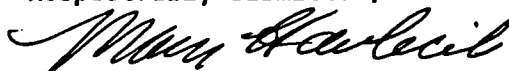
SAMPLE DESCRIPTION:
3E-9, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	14.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 21%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

MSD #5
E7563pc2.wr1/35
MH/bl/sc/sc

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San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7563
Collected: 08/05/88
Received: 08/07/88
Tested: 08/11 & 24/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3E-9, Soil
Duplicate Analysis of Same Extract

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ERM-WEST
WALNUT CREEK, CA

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	17.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 21%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
08/26/88
E7563pc3.wr1/37
MH/als/sc/sc

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Central Coast
Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7562
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

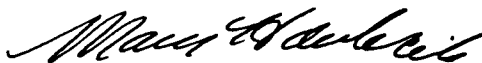
SAMPLE DESCRIPTION:
3E-11, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 17%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7562pc.wr1/35
MH/bl/sc/sc

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(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7561
Collected: 08/05/88
Received: 08/07/88
Tested: 08/11/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

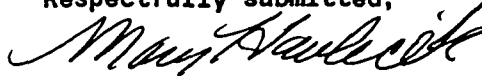
SAMPLE DESCRIPTION:
3E-13, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	3.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 23%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7561pc.wr1/35
MH/b1/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7560
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3E-15, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 13%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7560pc.wr1/35
MH/bl/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7694
Collected: 08/08/88 @ 1713
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596


SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3E-17 (2.0 to 2.5 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	1.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 13%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

ECD
E7694pc.wr1/36
MH/bl/sc/sc

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141 Suburban Road , Suite C-4
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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7685
Collected: 08/08/88 @ 1420
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3F-4 (2.0 to 2.5 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 8.0%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

ECD
E7685pc.wr1/36
MH/jg/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7559
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3F-6, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	7.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 20%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7559pc.wr1/35
MH/bl/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7558
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

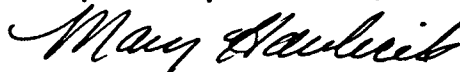
SAMPLE DESCRIPTION:
3F-8, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	4.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 19%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7558pc.wr1/34
MH/jc/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7557
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

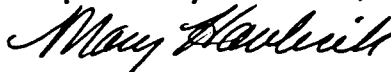
SAMPLE DESCRIPTION:
3F-10, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	2.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 14%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7557pc.wr1/34
MH/jc/sc/sc

Central
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Analytical
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Central Coast
Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7556
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3F-12, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 19%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7556pc.wr1/35
MH/bl/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7555
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

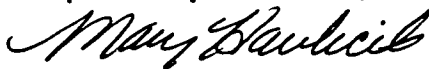
SAMPLE DESCRIPTION:
3F-14, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 17%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7555pc.wr1/35
MH/bl/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7693
Collected: 08/08/88 @ 1657
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

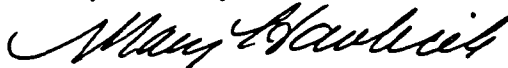
SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3F-16 (2.0 to 2.5 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 16%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

ECD
E7693pc.wr1/36
MH/bl/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7686
Collected: 08/08/88 @ 1435
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3G-5 (2.5 to 3.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 11%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

ECD
E7686pc.wr1/36
MH/jg/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7687
Collected: 08/08/88 @ 1451
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3G-7 (2.5 to 3.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 13%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

ECD
E7687pc.wr1/36
MH/jg/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7552
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3G-9, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	9.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 23%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7552pc.wr1/35
MH/bl/gh/gh

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7553
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3G-11, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	2.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 18%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7553pc.wr1/34
MH/jc/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7554
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3G-13, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	2.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 22%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
E7554pc.wr1/34
MH/jc/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7695
Collected: 08/08/88 @ 1605
Received: 08/09/88 @ 1800
Tested: 08/12/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3G-15 (1.5 to 2.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 5.3%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
E7695pc.wr1/36
MH/bl/sc/sc

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Lab Number: E-7688
Collected: 08/08/88 @ 1506
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
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Suite 260
Walnut Creek, CA 94596

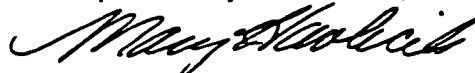
SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3H-6 (2.0 to 2.5 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 7.1%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

ECD
E7688pc.wr1/36
MH/jc/sc/sc

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San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7689
Collected: 08/08/88 @ 1520
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3H-8 (2.0 to 2.5 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 17%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek

Mary Havlicek, Ph.D.
President

ECD
E7689pc.wr1/36
MH/jc/sc/sc

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Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7551
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

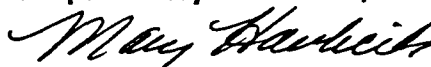
SAMPLE DESCRIPTION:
3H-10, Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 15%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7551pc.wr1/35
MH/bl/sc/sc

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141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7696
Collected: 08/08/88 @ 1625
Received: 08/09/88 @ 1800
Tested: 08/12/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

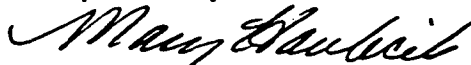
SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3H-12 (1.5 to 2.5 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 9.0%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7696pc.wr1/36
MH/bl/sc/sc

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Analytical Services
141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7690
Collected: 08/08/88 @ 1600
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3J-9 (2.5 to 3.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 11%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek

Mary Havlicek, Ph.D.
President

ECD
E7690pc.wr1/36
MH/jc/sc/sc

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Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7550
Collected: 08/05/88
Received: 08/07/88
Tested: 08/09/88
Collected by: A. Chemburkar

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
3J-11, Soil

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Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	2.

Compounds listed as "not found" would have been reported if present at or above the listed detection limits. Sample was extracted 08/08/88.

Percent Moisture: 17%. Tested by EPA Method 160.3 on 08/08/88 by ACF.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
E7550pc.wr1/34
MH/jc/sc/sc

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Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7697
Collected: 08/08/88 @ 1640
Received: 08/09/88 @ 1800
Tested: 08/12/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

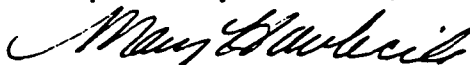
SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3J-13 (1.5 to 2.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 19%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
E7697pc.wr1/36
MH/bl/sc/sc

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141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7698
Collected: 08/08/88 @ 1702
Received: 08/09/88 @ 1800
Tested: 08/12/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3H-14 (1.5 to 2.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	1.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 12%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
E7698pc.wr1/36
MH/bl/sc/sc

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(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7691
Collected: 08/08/88 @ 1620
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3K-10 (2.5 to 3.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 8.9%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

ECD
E7691pc.wr1/36
MH/jc/sc/sc

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EPA METHOD 608/8080 - PCB'S

Lab Number: E-7692
Collected: 08/08/88 @ 1645
Received: 08/09/88 @ 1800
Tested: 08/11/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

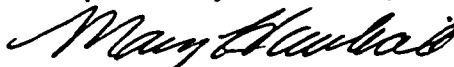
SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3K-12 (2.5 to 3.0 Feet), Soil

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	18.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 8.4%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

ECD
E7692pc.wr1/36
MH/jc/sc/sc

Central
Coast
Analytical
Services

Central Coast
Analytical Services
141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: E-7692
Collected: 08/08/88 @ 1645
Received: 08/09/88 @ 1800
Tested: 08/11 & 24/88
Collected by: CS/AC

ERM-WEST
1777 Botelho Dr.
Suite 260
Walnut Creek, CA 94596

SAMPLE DESCRIPTION:
Job #40058, Hunter's Point S.F.,
3K-12 (2.5 to 3.0 Feet), Soil
Duplicate Analysis of Same Extract

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	17.

Compounds listed as "not found" would have been reported if present at or above the listed detection limits. Sample was extracted 08/10/88.

Percent Moisture: 8.4%. Tested by EPA method 160.3 on 08/15/88 by PD.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

ECD
08/26/88
E7692pc2.wr1/37
MH/als/sc/sc

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Analytical
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Central Coast
Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: B-08118
Collected:
Received:
Tested: 08/11/88
Collected by:

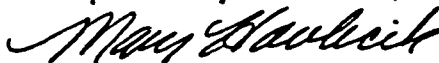
CCAS

SAMPLE DESCRIPTION:
Instrument Blank

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
B08118pc.wr1/35
MH/bl/sc/sc

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Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: QH-08108
Collected: 08/10/88
Received: 08/10/88
Tested: 08/11/88
Collected by:

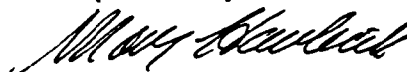
CCAS

SAMPLE DESCRIPTION:
Sonic Horn Extraction Blank
08/10/88

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/10/88.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
QH0810pc.wr1/35
MH/bl/sc/sc

Central
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Central Coast
Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: QS-08068-1
Collected: 08/06/88
Received: 08/06/88
Tested: 08/12/88
Collected by:

CCAS

SAMPLE DESCRIPTION:
Sand Spiked With Arochlor 1248

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.01	not found
PCB 1221	0.01	not found
PCB 1232	0.01	not found
PCB 1242	0.01	not found
PCB 1248	0.01	0.1 *
PCB 1254	0.01	not found
PCB 1260	0.01	not found

Percent Recovery of Arochlor 1248 = 100%.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/06/88.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
QS806pc1.wr1/36
MH/bl/sc/sc

Central
Coast
Analytical
Services

Central Coast
Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: QS-08068-3
Collected: 08/06/88
Received: 08/06/88
Tested: 08/12/88
Collected by:

CCAS

SAMPLE DESCRIPTION:
Sand Spiked With Arochlor 1248

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	0.01	not found
PCB 1221	0.01	not found
PCB 1232	0.01	not found
PCB 1242	0.01	not found
PCB 1248	0.01	0.07 *
PCB 1254	0.01	not found
PCB 1260	0.01	not found

Percent Recovery of Arochlor 1248 = 70%.

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/06/88.

Respectfully submitted,

Mary Havlicek
Mary Havlicek, Ph.D.
President

MSD #5
QS806pc3.wr1/36
MH/bl/sc/sc

Central
Coast
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Services

Central Coast
Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: B-08128
Collected:
Received:
Tested: 08/12/88
Collected by:

CCAS

SAMPLE DESCRIPTION:
Instrument Blank

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits.

Respectfully submitted,



Mary Havlicek, Ph.D.
President

MSD #5
B08128pc.wr1/36
MH/bl/sc/sc

Central
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Services

Central Coast
Analytical Services
141 Suburban Road , Suite C-4
San Luis Obispo, California 93401
(805) 543-2553
EPA METHOD 608/8080 - PCB'S

Lab Number: QH-08068
Collected: 08/06/88
Received: 08/06/88
Tested: 08/12/88
Collected by:

CCAS

SAMPLE DESCRIPTION:
Sonic Horn Extraction Blank
08/06/88

Compound Analyzed	Detection Limit milligrams/Kg	Concentration milligrams/Kg
PCB 1016	1.	not found
PCB 1221	1.	not found
PCB 1232	1.	not found
PCB 1242	1.	not found
PCB 1248	1.	not found
PCB 1254	1.	not found
PCB 1260	1.	not found

Compounds listed as "not found" would have been reported if present
at or above the listed detection limits. Sample was extracted 08/06/88.

Respectfully submitted,


Mary Havlicek, Ph.D.
President

MSD #5
QH0806pc.wr1/36
MH/bl/sc/sc

APPENDIX E

CHAIN OF CUSTODY FORMS
FOR VERIFICATION SAMPLING PROGRAMS

E-1 AREA 1 (Interim Report)
E-2 AREA 2
E-3 AREA 3

E-1 AREA 1 (Interim Report)

Security Seal: intact

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler (s): D. C. C. V. G. N. O.
Date: 4-15-87
Weather: Sunny, breezy

Job Location: Hunters Pt., San Francisco
Job No: 400-30
No. of Samples Collected: 21 soils
page 1 of 4

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification samples: 3 day turn around

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contrr. Type	Preservative	Iced (Y/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
D2680 A-1*	4:00P		X	602.	1 glass jar	Done	YES	trowel	See note below
2681 A-2	4:05P		X						
2682 B-1	4:10P		X						
2683 2 ft. S.E. of B-2	4:14P		X						
2684 C-1*	4:19P		X						
2685 C-2	4:24P		X						

* split sample to DONS

Comments:

Verification samples
for PCB cleanup

Composite the following
sample groups for
single discrete analyses (PCBs)

- A-1, A-2, B-2, B-1
- C-1, C-2, D-1, D-2, D-3

Custody Record

Signature, Date/Time
Relinquished: D. C. C. V. G. N. O. 4-16-87 9 AM
Received: 12-11-87 1720H
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory

Central Coast
Analytical Services
141 Shubert Rd
San Luis Obispo, CA
93401
(805) 543-2553

Security Seal: intact 13

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler (s): D. C. C. C.
Date: 4-15-87
Weather: sunny, hazy, warm

Job Location: Hunter's Pt., San Francisco CA
Job No: 400-30
No. of Samples Collected: 21 soils
page 2 of 4

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification samples: 3 day turn around

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contr. Type	Preservative	Iced (Y/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
D2686 D-1	4:28P		X	602-	1 glass jar	none	YES	trowl	see note below
2687 D-2*	4:34P		X						
2688 D-3	4:38P		X						
2689 E-1	4:42P		X						
2690 E-2	4:45P		X						
2691 E-3	4:49P		X						

* split sample to D045

Comments:

Verification samples
composite the following
sample apps. for single
discrete analyses (PDB)
(3) E-1, E-2, E-3 F-1
(4) F-2, F-3, G-1, G-2

Custody Record

Signature, Date/Time
Relinquished: D. C. C. C. 4-16-87 9 AM
Received: D. C. C. C. 4-16-87 12:00 PM
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory

Central Coast
Analytical
San Luis Obispo CA

Security Seal: intact

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler (s): D. Cataggo
Date: 4-15-87
Weather: sunny, breezy, warm

Job Location: Hunters Pt., San Fran CA
Job No: 440-30
No. of Samples Collected: 21 soils
page 3 of 4

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification samples: 3 day turn around

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contr. Type	Preservative	Iced (Y/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
02692 F-1*	4:55P		X	6002.	1 glass; 3-5	None	YES	4 Gwells	see note below
2693 F-2	5:00P		X						
2694 F-3	5:04P		X						
2695 3.5 ft. N. of G-1	5:09P		X						
2696 G-2	5:14P		X						
2697 G-4	5:19P		X						

* split sample to DOTS

Comments:

Verification samples

comparing the following sample sets for single discrete analyses (PCBs)

(4) F-2, F-3, G-1, G-2
(5) G-4, H-1, H-2, H-3

Custody Record

Signature, Date/Time

Relinquished: D. Cataggo 4-16-87 9 AM

Received: [Signature]

Relinquished: _____

Received: _____

Relinquished: _____

Received: _____

Relinquished: _____

Received: _____

Name and Address of Receiving Laboratory

Central Coast Analytical

San Luis Obispo CA

Security Seal: intact!!

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler (s): D. Cutugno
Date: 4-15-87
Weather: Sunny, breezy, warm

Job Location: Avalon Pt., San Fran CA
Job No: 400-30
No. of Samples Collected: 21 Soils
page 4 of 4

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification samples: 3 day turn around

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contrn. Type	Preservative	Iced (Y/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
D2698 H-2	S:24 P		X	6 oz.	1 glass jar	None	YES	Acid	See note below
2699 H-3*	S:30 P		X	↓	↓	↓	↓	↓	↓
2700 H-1	S:37 P		X	↓	↓	↓	↓	↓	↓

Comments: * split sample to DOHS
Verification samples
composited the following
sample grps. for single
discrete analyses (PEB):
(S) G-4, H-1, H-2, H-3

Custody Record
Signature, Date/Time
Relinquished: D. Cutugno 4-16-87 9 AM
Received: 17/1/87 / CC/PS 1700 4/16
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory
Central Coast
Analytical
San Luis Obispo CA

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: NAVY
Sampler(s): MILLER/BROWN
Date: 5/18/87
Weather: Foggy/overcast A.M.

Job Location: Hunters Point
Job No: 40030
No. of Samples Collected: 16
page 1 of 3

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

3 day Turn Around

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contrr. Type	Preservative	Iced (Y/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
X1	4:55 PM		X	3-403	1/ GLASS	None	Y	Auger (hand) Traveller to Container	PCB
X2	4:45 PM		X						
X3	4:32 PM		X						
X4	4:25 PM		X						
X5	5:05 PM		X						
X6	5:12 PM		X						
X7	5:20 PM		X						

Comments:

SAMPLE DEPTHS FROM SURFACE
X1 = 2'-6"
X2 = 2'-2"
X3 = 2'-4"
X4 = 2'-0"
X5 = 2'-0"
X6 = 2'-2"
X7 = 1'-11"

Custody Record

Signature, Date/Time
Relinquished: Dennis Miller 5-18-87
Received: 7:00 PM
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory

Central Coast Analytical
141 Suburban Rd Suite C-4
San Luis Obispo, CA 93401
805-543-2553

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Client: NAVY
Sampler (s): MILNER/BROWN
Date: 5-18-87
Weather: Foggy/overcast A.M. / Sunny, windy P.M.

Job Location: HUNTERS Point
Job No: 40030
No. of Samples Collected: 16
page 1 of 3

3 day Turn Around

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contrn. Type	Preservative	Iced (X/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
X 8	5:15 PM		X	~3403	1/GLASS	NONE	Y	AUGER (HAND) Trowelled to Container	PCB
X 9	5:07 PM		X	}	}	}	}		}
X 10	5:01 PM		X	}	}	}	}		}
X 11	4:27 PM		X	}	}	}	}		}
X 12	3:45 PM		X	}	}	}	}		}
X 13	3:55 PM		X	}	}	}	}		}
X 14	4:02 P.M.		X	}	}	}	}		}

Comments:

Sample depths from Surface
X 8 = 1'-10"
X 9 = 2'-1"
X 10 = 1'-9"
X 11 = 2'-0"
X 12 = 2'-4"
X 13 = 2'-0"
X 14 = 2'-0"

Custody Record

Signature, Date/Time
Relinquished: Dennis Miller 5-18-87
Received: 7:00 PM
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: Phyllis Miller
Received: _____

Name and Address of Receiving Laboratory

Central Coast Analytical
141 Suburban Rd. Suite C-4
San Luis Obispo, CA 93401
805-543-2553

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: NAVY Job Location: Hunters Point
 Sampler (s): MILLER/BROWN Job No: 40030
 Date: 5/18/87 No. of Samples Collected: 16
 Weather: Foggy / overcast A.M. / Sunny / Windy P.M.
 page 3 of 3

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

3 day Turn Around

Sample ID #	Time	Sample Type		Volume	No. of Contnrs. Contnr. Type	Preservative	Iced (X/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
X15	4:10 PM		X	3-403	1 GLASS	None	Y	Auger (HAND)	PCB
X16	4:15 PM		X	"	↓	None	Y	1 rowelled to container	PCB

Comments:

Sample depths from surface:

X15 = 2'-0"
X16 = 2'-6"

Custody Record

Signature, Date/Time
 Relinquished: Dennis Miller 5-18-87
 Received: 7:00 PM
 Relinquished: _____
 Received: _____
 Relinquished: _____
 Received: _____
 Relinquished: Therrell W. Miller

Name and Address of Receiving Laboratory

Central Coast Analytical
141 Suburban Rd. Suite C-4
San Luis Obispo, CA 93401
805-543-2553

E-2 AREA 2

HAZARDOUS MATERIALS SAMPLE ANALYSIS REQUEST		All applicable items must be completed		1. HML No. To: <u>D500-D502</u>		2. Page <u>1</u>	
3. Collector <u>chein kao</u>		4. Phone (818) <u>415 540</u> 557 <u>3052</u>		5. Priority <u>(2)</u>		6. Authorized by _____	
7. Date Sampled <u>9/3/87</u>		7. Time Sampled <u>11:00</u> Hours		8. Codes (fill in all applicable codes)			
9. Activity <input type="checkbox"/> Ent <input type="checkbox"/> Surv <input checked="" type="checkbox"/> Site Mit <input type="checkbox"/> Permitting <input type="checkbox"/> Alt Tech <input type="checkbox"/> Other				a. STC <u>001021</u>			
10. SAMPLING LOCATION				b. Region <u>4</u>			
a. EPA ID No.				c. TPC _____			
b. Site <u>Hunters Point Naval Shipyard</u>				d. INDEX <u>6530</u>			
c. Address <u>S. F.</u>				e. PCA <u>11025</u>			
Number _____ Street _____ City _____ Zip _____				f. SITE <u>2000500</u>			
				g. County <u>075</u>			
11. SAMPLES							
a. ID	b. Collector's No.	c. HML No.	d. Type	e. Type	f. Size	g. Field Information	
A. <u>D2392A-1</u>		<u>D500</u>	<u>soil</u>	<u>G</u>	<u>250</u>	<u>Soil boring at 1'11" to 2'6"</u>	
B. <u>40 2E-1</u>		<u>D501</u>	<u>soil</u>	<u>G</u>	<u>250</u>	<u>Soil boring at 11" to 1'14"</u>	
C. <u>4 2C-6</u>		<u>D502</u>	<u>soil</u>	<u>G</u>	<u>250</u>	<u>Soil boring at 1' to 1'11"</u>	
D.							
E.							
F.							
G.							
H.							
12. ANALYSIS REQUESTED							
a. <input type="checkbox"/> pH		f. <input checked="" type="checkbox"/> PCB		k. <input type="checkbox"/> Ext. Org (Screening)			
b. <input type="checkbox"/> Metal Scan		g. <input type="checkbox"/> VOA		l. <input type="checkbox"/> Chlorinated Pesticides			
c. <input type="checkbox"/> Metals (Spec)		h. <input type="checkbox"/> PAH		m. <input type="checkbox"/> Organo-P Pesticides			
d. <input type="checkbox"/> W.E.T.		i. <input type="checkbox"/> Phenols		n. <input type="checkbox"/>			
		j. <input type="checkbox"/> Carba-mates		o. <input type="checkbox"/>			
13. CHAIN OF CUSTODY							
a.	<u>chein kao</u>	<u>chein ping kao / AWME</u>	<u>9/3/87 - 9/13/87</u>				
b.	<u>[Signature]</u>	<u>VERDE EARL WOODS / LAB ASST</u>	<u>9/13/87 - 9/15/87</u>				
c.	<u>chein kao</u>	<u>chein ping kao / AWME</u>	<u>9/15/87 - 9/16/87</u>				
d.	<u>Shellie Noller</u>	<u>Shellie Noller / Sample Control</u>	<u>9/17/87 - 1/1</u>				
Security seal intact upon receipt							
HML to prep sample for split for inspection							
14. SPECIAL REMARKS							
RECEIVED BY <u>[Signature]</u> a. Title <u>PHC III</u> b. Date <u>9-3-87</u>							
15. SAMPLE ALLOCATION a. <input type="checkbox"/> HML-Berkeley b. <input type="checkbox"/> HML-SC c. <input type="checkbox"/> AIHL d. <input type="checkbox"/> Contract b. Date _____							
16. ANALYSIS REQUESTED <u>PCB</u>							

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler(s): D. Cufugno
Date: 9-3-87
Weather: Sunny, breezy, warm, dry

Job Location: San Fran CA
Job No: 400-36
No. of Samples Collected: 14 soils

page 1 of 3

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification Samples: 5-day Turn around

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contrr. Type	Preservative	Iced (X/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
✓ 2A-1 16"-2'0"	9:50A		D-6886 X	2" x 6"	1 brass tube	none	YES	shive sampler	PCBs
✓ 2G-1 18"-2'3"	10:15A		6887 X	2" x 7"	1 steel tube				
✓ 2F-1 15"-2'0"	10:45A		6888 X	2" x 7"	"				
✓ 2D-0 22"-2'8"	11:45A		6889 X	2" x 6"	1 brass tube				
✓ 2B-6 16"-2'0"	12:55P		6890 X	2" x 6"	"				
✓ 2E-7 27"-3'1"	2:25P		6891 X	2" x 6"	"				

Comments: Analyze soil
from bottom (re-use for
depth) of each hole
Soil, fines & gravel: ERM
NOT INCLUDE ROCKS
CONTAMINATE ERM.
Analyze 1 random sample
in duplicate (1 of 14 total)

Custody Record
Signature, Date/Time
Relinquished: D. Cufugno 9-3-87 6:45P
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory
Central Coast
Analytical Services (CCAS)
141 Suburban Rd
S-4
San Luis Obispo CA
93401
(805) 393-2553

NOTE: Samples 2A-1, 2C-6 and 2E-1 were retained by the Dept. of Health Services. Splits of these samples will be sent to ERMS under separate cover.

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler(s): Dr. Cufuano
Date: 9-3-87
Weather: see p. 1

Job Location: San Fran CA
Job No: 400-36
No. of Samples Collected: 14 soils
page 2 of 3

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification Samples: 5-day turn around

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contr. Type	Preservative	Iced (Y/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab Comp.						
✓ 2D-6 16"-2'0"	2:35P		1) (6.8.87) X	2" x 6"	1 steel tube	none	YES	drive sampler	PCBs
✓ 2E-7 18"-2'2"	2:50P		(6.8.87) X		1 brass tube				
✓ 2F-7 1'10"-2'4"	3:10P		(6.8.87) X		"				
✓ 2G-6 2'4"-2'10"	3:25P		(6.8.87) X		"				
✓ 2H-6 1'4"-1'10"	3:50P		(6.8.87) X		1 steel tube				
✓ 2B-1 18"-2'2"	4:10P		(6.8.87) X		"				

Comments: _____
_____ see correct section
_____ p. 1

Custody Record
Signature, Date/Time
Relinquished: Dr. Cufuano 9-3-87 6:45P
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: Whittier, Whittier
Received: 9-4-87 0900

Name and Address of Receiving Laboratory

Central Coast
Analytical Services

San Luis Obispo CA

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler (s): D. Cutyne
Date: 9-3-87
Weather: see p-1

Job Location: San Fran CA
Job No: 400-36
No. of Samples Collected: 14 soils

page 3 of 3

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification Samples: 5-day turn around

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contr. Type	Preservative	Iced (X/N)	Sampling Method	Analyses	
		Water Comp.	Soil Grab							
2C-1 1'6"-2'0"	4:20P	1)	6.8"	X	2" X 6"	1 Grass tube	none	YES	drive sampler	PCBs
BL-1 1'6"-2'0"	4:35P		6.8"	X	"	"	"	"	"	"

Comments: _____
see comment section
pg 1.

Custody Record
Signature, Date/Time
Relinquished: D. Cutyne 9-3-87 6:45P
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: Walter Miller
Received: 9-4-87 0900

Name and Address of Receiving Laboratory

Central Coast
Analytical Services

San Luis Obispo CA

CHAIN OF CUSTODY SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy Job Location: San Fran, CA
 Sampler (s): D. Catugno/R. Kuyf Job No: 400-36
 Date: 9/4/87 No. of Samples Collected: 31 soils / 2 waters
 Weather: partly cloudy, warm, breezy, dry
 page 1 of 16

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification Samples : 5-day turnaround

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contr. Type	Preservative	Iced (X/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab Comp. Grab						
6950 2A-2	2:13 P		X	~4 oz.	1 glass jar	none	YES	manual	PCBs
6951 2A-3	2:17 P		X						
6952 2A-4	2:22 P		X						
6953 2B-2	2:20 P		X						
6954 2B-3	2:18 P		X						
6955 2B-4	2:15 P		X						

Comments:

31 ~~soils~~ soils : analyze
 10% to duplicate

Custody Record
 Signature, Date/Time
 Relinquished: D. Catugno 9/4/87 6PM
 Received: _____
 Relinquished: _____
 Received: _____
 Relinquished: _____
 Received: _____
 Relinquished: _____
 Received: _____

Name and Address of Receiving Laboratory
Central Coast
Analytical Services
141 Suburban Rd
C-4
San Luis Obispo CA
93401
(805) 543-2553

NOTE: Samples 2C-2, 2C-5, 2E-4 and 2G-4 were retained by the Dept. of Health Services; splits will be sent to CCAS

CHAIN OF CUSTODY & SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler(s): Cutugno/Krapp
Date: 9/4/87
Weather: see p. 1

Job Location: San Francisco
Job No: 400-36
No. of Samples Collected: 3:1 soils/water

page 2 of 6

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification Samples: 5-day turnaround

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contrn. Type	Preservative	Iced (Y/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
0956 2B-5	2:28P		X	~4 oz.	1 glass jar	none	YES	manual	PCBs
0957 2C-3	2:24P		X						
0958 2C-4	2:24P		X						
0959 2D-1	2:36P		X						
0960 2D-2	2:39P		X						
0961 2D-3	2:35P		X						

Comments: _____
_____ see p. 1 _____

Custody Record
Signature, Date/Time
Relinquished: D. Cutugno 9/4/87 CPM
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory

Central Coast
Analytical
San Luis Obispo CA

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler(s): Cutugno/Knap
Date: 9/4/87
Weather: see p. 1

Job Location: San Fran CA
Job No: MOD-36
No. of Samples Collected: 31 water/2 soils

page 3 of 6

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification Samples: 5-day turnaround

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contrr. Type	Preservative	Iced (X/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
961 2D-4	2:43 P		X	~4 oz.	1 Glass	none	YES	manual	PCBs
963 2D-5	2:12 P		X						
964 2E-2	2:48 P		X						
965 2E-3	2:49 P		X						
966 2E-5	2:56 P		X						
967 2E-6	3:10 P		X						

Comments: _____

see p. 1

Custody Record
Signature, Date/Time
Relinquished: D. Cutugno 9/4/87 GPM
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory

Central Coast
Analytical
San Luis Obispo

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler(s): C. Tuono/Knapp
Date: 9/4/87
Weather: see p. 1

Job Location: San Fran CA
Job No: WOD-36
No. of Samples Collected: 36 soils/2 waters

page 4 of 6

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification Samples: 5-day turnaround

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contrr. Type	Preservative	Iced (X/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
D6468 2F-2	2:50 P		X	~402	1 glass jar	none	YES	manual	PCBs
6969 2F-3	2:53 P		X						
6970 2F-4	3:31 P		X						
6971 2F-5	3:20 P		X						
6972 2F-6	3:35 P		X						
973 2G-2	3:01 P		X						

Comments: _____

see p. 1

Custody Record
Signature, Date/Time
Relinquished: H. Tuono 9/4/87 CPM
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory

Central Coast

Analytical Services

San Luis Obispo CA

CHAIN OF CUSTODY AND SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Navy
Sampler(s): C. A. Lugo / K. W. P.
Date: 9/4/97
Weather: see p. 1

Job Location: San Francisco
Job No: 400-36
No. of Samples Collected: 31 soils / 2 waters

page 3 of 5

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification Samples: 5-day turnaround

Sample ID #	Time	Sample Type		Volume	No. of Contnrs. Contrn. Type	Preservative	Iced (X/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
6974 2G-3	2:55P		X	~402.	1 glass jar	none	YES	manual	PCBs
6975 2G-5	3:23P		X						
6976 2H-2	3:02P		X						
6977 2H-3	3:14P		X						
6978 2H-4	3:17P		X						
6979 2H-5	3:43P		X						
6980 BL-2	5:00P		X	~202.	1 glass jar	none	YES	manual	PCBs

Comments: _____

see p. 1

Custody Record
Signature, Date/Time
Relinquished: D. Cantuano 9/4/97 6PM
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory

Central Coast
Analytical

San Luis Obispo CA

CHAIN OF CUSTODY SAMPLE IDENTIFICATION RECORD

ERM-West
Environmental
Resources
Management

Client: Nav.
Sampler(s): C. H. Jones
Date: 9/4/87
Weather: see p. 1

Job Location: San Francisco
Job No: 400-36
No. of Samples Collected: 31 soils/2 waters

page 6 of 6

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

5-day turnaround

D6981
D6982

Sample ID #	Time	Sample Type		Volume	No. of Contnrs. Contnr. Type	Preservative	Iced (X/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab Comp. Grab						
#1	9:45A		X	12	1 amber glass bottle	None	YES	manual	see comment section
#2	3:20P		X	12	"	"	"	"	"

Comments: _____

PCB Analysis:
#1: supernatant
only: DO NOT include
SEDIMENT
#2: Total contents

Custody Record
Signature, Date/Time
Relinquished: D. C. Jones 9/4/87 6PM
Received: [Signature]
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory

Central Coast
Analytical
San Luis Obispo CA

CHAIN OF CUSTODY & SAMPLE IDENTIFICATION RECORD

ERK-West
Environmental
Resources
Management

Client: Navy
Sampler (s): D. Cutugno
Date: 10-16-87
Weather: sunny, warm, dry

Job Location: Hunters Pt., San Fran.
Job No: 400-36
No. of Samples Collected: 11*

1777 Botalbo Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0435

page 1 of 2

Verification Samples: 5-day turnaround

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contr. Type	Preservative	Iced (X/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
2C-5a	12:31P		X	N6oz.	1 glass jar	none	YES	manual	see comment section
2C-5b	12:32P		X						
2C-5c	12:34P		X						
2E-4	12:36P		X						
2D-2	12:37P		X						
2E-3b	12:39		X						

Comments: _____

Analyze for PCBs;
if possible, ATR DAY
prior to screening
and homogenizing
Analyze 2E-2 in
TRIPLICATE

Custody Record
Signature, Date/Time
Relinquished: D. Cutugno 10-16-87
Received: 2:45P
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory
Central Coast
Analytical Services
141 Suburban Rd
C-4
San Luis Obispo CA
93401
(805) 543-2553

* **NOTE:** Samples 2E-3 and 2C-5 were retained by CA.
DHS and will be sent to CCAS under separate
cover.

CHAIN OF CUSTODY D SAMPLE IDENTIFICATION RECORD

ERK-West
Environmental
Resources
Management

Client: Navy
Sampler(s): D. Cutugno
Date: 10-16-87
Weather: swampy, warm, dry

Job Location: Hunter Pt., San Fran.
Job No: 400-36
No. of Samples Collected: 11* (see p.1)

page 2 of 2

1777 Botelho Drive
Suite 260
Walnut Creek, CA 94596
(415) 946-0455

Verification Samples: 5-day turnaround

Sample ID #	Time	Sample Type		Volume	No. of Contrs. Contrr. Type	Preservative	Iced (Y/N)	Sampling Method	Analyses
		Water Comp.	Soil Grab						
2E-3a	12:42P		X	~6oz.	1 glass jar	none	YES	manual	see comment section pg 1
2E-2a	12:45P		X	↓	↓	↓	↓	↓	↓
2E-2	1:05P		X	↓	↓	↓	↓	↓	↓

Comments: _____

Custody Record
Signature, Date/Time
Relinquished: D. Cutugno 10-16-87
Received: 2:45P
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____
Relinquished: _____
Received: _____

Name and Address of Receiving Laboratory

Central Coast
Analytical Svcs.
San Luis Obispo CA

HAZARDOUS MATERIALS
SAMPLE ANALYSIS REQUESTAll applicable items
must be completed1. HML No. To D723-D724 2. Page of3. Collector Chein Kao4. Phone 415 540-30525. Priority ☐
a. Authorized by _____6. Date Sampled 10/16/877. Time Sampled 12.30 Hours

8. Codes (fill in all applicable codes)

9. Activity ☐ Ent ☐ Surv ☒ Site Mit ☐ Permitting ☐ All Tech ☐ Other

10. SAMPLING LOCATION

a. EPA ID No.

b. Site Hunters Point Naval Shipyardc. Address S.F.

Number

Street

City

Zip

a. STC

b. Region

c. TPC

d. INDEX

e. PCA

f. SITE

g. County

11. SAMPLES

Container

g. Field Information

a. ID

b. Collector's No.

c. HML No.

d. Type

e. Type

f. Size

g. Field Information

A

2C-5

D723

SOIL

G

250

resample the same loc

B

2E-3

D724

SOIL

G

250

"

C

D

E

F

G

H

12. ANALYSIS REQUESTED

1. ☒ PCBk. ☐ Ext Org
(Screening)a. ☐ pHg. ☐ VOAl. ☐ Chlorinated
Pesticidesb. ☐ Metal
Scanh. ☐ PAHm. ☐ Organo-P
Pesticidesc. ☐ Metals
(Spec)i. ☐ Phenolsn. ☐d. ☐ W.E.T.j. ☐ Carba-
mateso. ☐

13. CHAIN OF CUSTODY

a. Chein Ping Kao

Signature

Chein Ping Kao/AWME

Name/Title

10/16/87-10/16/87

Inclusive Dates

b. VERUE EARL WORKS/LABASIT

Signature

VERUE EARL WORKS/LABASIT

Name/Title

10/16/87-10/23/87

Inclusive Dates

c. Chein Ping Kao

Signature

Chein Ping Kao/AWME

Name/Title

10/23/87-11/1/87

Inclusive Dates

d. Shelli Miller

Signature

Shelli Miller/sample control - CCAS

Name/Title

10/26/87-11/1/87

Inclusive Dates

14. SPECIAL REMARKS

please contact Howard Okamoto for special instructions

15. RECEIVED BY

Howard S. Okamoto

a. Title

PHC III

b. Date

10/16/87

16. SAMPLE ALLOCATION

a. ☐ HML-Berkeleyb. ☐ HML-SCc. ☐ AIHLd. ☐ Contract

b. Date

17. ANALYSIS REQUESTED

PCBs

Air dried #10 mesh fraction - splits to be given to C.I. Kao

NLY (PCL)

HAZARDOUS MATERIALS SAMPLE ANALYSIS REQUEST		All applicable items must be completed		1. HML No. To: <u>DS09-D12</u>		2. Page 2 of 2	
Collector <u>Chein Kao</u>				4. Phone <u>AK 540-3052</u>			
6. Date Sampled <u>9/4/87</u>		7. Time Sampled		Hours		8. Codes (fill in all applicable codes)	
9. Activity <input type="checkbox"/> Ent <input type="checkbox"/> Surv <input checked="" type="checkbox"/> Site Mit <input type="checkbox"/> Permitting <input type="checkbox"/> Alt Tech <input type="checkbox"/> Other				5. Priority <input checked="" type="checkbox"/>			
10. SAMPLING LOCATION				a. Authorized by			
a. EPA ID No.				a. STC			
b. Site <u>Hunters Point Naval Shipyard</u>				b. Region <u>4</u>			
c. Address <u>S.F.</u>				c. TPC			
Number Street City Zip				d. INDEX <u>6530</u>			
				e. PCA <u>11025</u>			
				f. SITE <u>2000500</u>			
				g. County <u>075</u>			

11. SAMPLES						
a. ID	b. Collector's No.	c. HML No.	d. Type	e. Type	f. Size	g. Field Information
<u>1. DPM 2C-5</u>		<u>DS09</u>	<u>Soil</u>	<u>G</u>	<u>250</u>	
<u>2. 23 2B4</u>		<u>DS10</u>	<u>"</u>	<u>G</u>	<u>"</u>	
<u>3. 44 2G-4</u>		<u>DS11</u>	<u>"</u>	<u>G</u>	<u>"</u>	
<u>4. 45 2C-2</u>		<u>DS12</u>	<u>"</u>	<u>G</u>	<u>"</u>	
<u>E</u>						
<u>F</u>						
<u>H</u>						

12. ANALYSIS REQUESTED		
a. <input type="checkbox"/> pH	f. <input checked="" type="checkbox"/> PCB	k. <input type="checkbox"/> Ext. Org (Screening)
b. <input type="checkbox"/> Metal Scan	g. <input type="checkbox"/> VOA	l. <input type="checkbox"/> Chlorinated Pesticides
c. <input type="checkbox"/> Metals (Spec)	h. <input type="checkbox"/> PAH	m. <input type="checkbox"/> Organo-P Pesticides
d. <input type="checkbox"/> W.E.T.	i. <input type="checkbox"/> Phenols	n. <input type="checkbox"/>
	j. <input type="checkbox"/> Carbamates	o. <input type="checkbox"/>

13. CHAIN OF CUSTODY		
a. <u>Chein Kao</u> Signature	<u>Chein Ping Kao / AWME</u> Name/Title	<u>9/4/87 - 9/10/87</u> Inclusive Dates
b. <u>J. Thom Coons</u> Signature	<u>J. Thom Coons / Chemist</u> Name/Title	<u>9/14/87 - 9/14</u> Inclusive Dates
c. <u>Verne Earl Yabsco</u> Signature	<u>VERNE EARL YABSCO / HABSIT</u> Name/Title	<u>9/14/87 - 9/15</u> Inclusive Dates
d. <u>Chein Kao</u> Signature	<u>Chein Ping Kao / AWME</u> Name/Title	<u>9/15/87 - 9/16</u> Inclusive Dates
e. <u>Shellie Noller</u> Signature	<u>Shellie Noller / Sample Control</u> Name/Title	<u>9/17/87</u> Inclusive Dates

14. SPECIAL REMARKS <u>Security seal intact upon receipt.</u>	
15. RECEIVED BY <u>[Signature]</u>	16. Date <u>9-8-87</u>
17. SAMPLE ALLOCATION a. <input type="checkbox"/> HUAL Berkeley b. <input type="checkbox"/> HML-SC c. <input type="checkbox"/> AIHL d. <input type="checkbox"/> Contract	
18. ANALYSIS REQUESTED <u>PCB</u>	

E-3 AREA 3

ERM-West

1777 Botelho Drive • Suite 260 • Walnut Creek, CA • 94596 • (415) 946-0455

Date 8.5.88 Weather cloudy 65-70°F Page 1 of 3

Chain of Custody Record

Job # 40058					Collection			GC				GC/MS		Inorg	Other	Number of Containers	Remarks <u>5-Day Turnaround</u> <u>EPA Protocol</u>							
Job Location <u>HUNTER'S PT.</u>					Container type	ICED	Preservative	Sampling method	TPH-Extraction	BTEX/Total Fuel HCs	601 / 8010 Halocarbons	602 / 8020 Aromatics	604 / 8040 Phenols	8080 Pest/PCBs	624-8240 Purgeables			625-8270 BNAs & Pest (SVs)	Dioxins	Metals	Wet Chemistry	% moisture		
Sampler (signature) <u>ASC / Don Lapin</u>																								
Printed name <u>ARUNCHEMBURKAR / Don Lapin</u>																								
Lab Report Recipient																								
Telephone No.																								
Receiving Lab <u>CENTRAL COAST ANA. SER.</u>																								
Address <u>141 SUBURBAN ROAD, SUITE G-4</u>																								
Address <u>SAN LUIS OBISPO, CA 93401</u>																								
Sample ID#	Time	W-water S-soil	C-comp G-grab	Volume																				
31-11	1124	S	G		glass jar	Y	N	E-755														1	PCB 8080	
34-10	1129							7551															1	} % Moisture
36-9	1136							7552															1	
36-11	1142							7553															1	
36-13	1147							7554															1	
3F-14	1151							7555															1	
3F-12	1157							7556															1	
3F-10	1202							7557															1	
3F-8	1206							7558															1	
3F-6	1212							7559															1	

Precautions:

Conc: ☒ Lo ☐ Med ☐ HI Ship Via

Total Number of Containers:

10

Sample Relinquished By	Date	Time	Received By	Date	Time	Reason for Transfer (List Shipping Bill Number)
Anur	08/05		<i>Melinda H. ...</i>	8/7	1234	
Company <u>ERM-W</u>			Company <u>CC-4</u>			
Company			Company			

LABORATORY— Please Complete

☐ Samples Intact
 ☐ Samples at 4°C
 ☐ Samples not leaking
 ☐ # of containers matches C-of-C
 ☐ Container tags match C-of-C
 ☐ Cooler seals Intact

Lab sample custodian Signature _____ Date _____ Time _____

Sample Disposition
 ☐ Return to Site
 ☐ Discard
 ☐ Hold _____ days

7/28/88 D.L.

ERM-West

1777 Botelho Drive • Suite 260 • Walnut Creek, CA • 94596 • (415) 946-0455

Date 8.5.88 Weather cloudy 65-70°F Page 1 of 3

Chain of Custody Record

Job # <u>40058</u>					Collection			GC					GC/MS		Inorg	Other	Number of Containers	Remarks		
Job Location <u>HUNTER'S PT.</u>					Container type	ICED	Preservative	Sampling method	TPH-Extraction	BTEX/Total Fuel HCs	601 / 8010 Halocarbons	602 / 8020 Aromatics	604 / 8040 Phenols	608 / 8080 Pest/PCBs	624-8240 Purgeables	625-8270 BNAs & Pest (SVs)			Dioxins	Metals
Sampler (signature) <u>ASC / Don Lapin</u>																				
Printed name <u>ARUNCHEMBURKAR / DON LAPIN</u>																				
Lab Report Recipient																				
Telephone No.																				
Receiving Lab <u>CENTRAL COAST ANA. SER.</u>																				
Address <u>141 SUBURBAN ROAD, SUITE G-4</u>																				EPA Protocol
<u>SAN LUIS OBISPO, CA 93401</u>																				
Sample ID#	Time	Wettest Seal	C-comp Grab	Volume																
3E-15	1217	S	G	Glass jar	Y	N	7560													PCB 8080
3E-13	1221						7561													% Moisture
3E-11	1223						7562													
3E-9	1228						7563													
3E-7	1231						7564													
3D-12	1238						7565													
3D-10	1241						7566													
3D-8	1246																			
3D-6	1251						7567													
3D-4 3C-7	1258						7568													
Precautions:					Conc: <input checked="" type="checkbox"/> Lo <input type="checkbox"/> Med <input type="checkbox"/> HI					Ship Via					Total Number of Containers: <u>9</u>					
Sample Relinquished By		Date	Time	Received By		Date	Time	Reason for Transfer (List Shipping Bill Number)												
Arun		08	05	Marta H. ...		8/7	1254	1254												
Company				Company																
Company				Company																
LABORATORY—		<input type="checkbox"/> Samples Intact		<input type="checkbox"/> Samples at 4°C		<input type="checkbox"/> Samples not leaking		<input type="checkbox"/> # of containers matches C-of-C		<input type="checkbox"/> Container tags match C-of-C		<input type="checkbox"/> Cooler seals Intact								
Please Complete		Signature		Date		Time		Sample Disposition		<input type="checkbox"/> Return to Site		<input type="checkbox"/> Discard		<input type="checkbox"/> Hold _____ days						

Chain of Custody Record

Job #						Collection			GC			GC/MS			Inorg	Other	Remarks					
Job Location						Container type	ICED	Preservative	Sampling method	TPH-Extraction	BTEX/Total Fuel HCs	601 / 8010 Halocarbons	602 / 8020 Aromatics	604 / 8040 Phenols	606 / 8080 Pes/PCBs	624-8240 Purgeables		625-8270 BNAs & Pest (SVs)	Dioxins	Metals	Wet Chemistry	Number of Containers
Sampler (signature) ASC /																						
Printed name ARUNCHEMBURAR /																						
Lab Report Recipient																						
Telephone No.																						
Receiving Lab CENTRAL COAST ANA. SER.																						
Address 141 SUBURBAN ROAD, SUITE G-4																						
SAN LUIS OBISPO, CA 93401																						
Sample ID#	Time	Weather Sunset	C-comp G-grab	Volume																		EPA Protocol
3D-8	1302	S	G	6	✓	N	7569															PCB 8080
3C-9	1305						7570															+ % Moisture
3C-11	1309						7571															
3B-8	1316						7572															
3B-10	1318						7573															
3B-12	1321						7574															
3C-13	1328						7575															
3C-15	1332						7576															
3D-14	1337						7577															
Precautions:						Conc: <input checked="" type="checkbox"/> Lo <input type="checkbox"/> Med <input type="checkbox"/> HI			Ship Via			Total Number of Containers:			9							
Sample Relinquished By		Date	Time	Received By		Date	Time	Reason for Transfer (List Shipping Bill Number)														
Arun		06/05		Marta Henriquez		8/8	18341															
Company				Company																		
Company				Company																		
LABORATORY— Please Complete																						
<input type="checkbox"/> Samples Intact		<input type="checkbox"/> Samples at 4°C		<input type="checkbox"/> Samples not leaking		<input type="checkbox"/> # of containers matches C-of-C		<input type="checkbox"/> Container tags match C-of-C		<input type="checkbox"/> Cooler seals Intact												
Signature		Date		Time		Sample Disposition		<input type="checkbox"/> Return to Site		<input type="checkbox"/> Discard		<input type="checkbox"/> Hold _____ days										

Chain of Custody Record

Job # <u>40058</u>					Collection			GC				GC/MS		Inorg	Other	Number of Containers	Remarks				
Job Location <u>Hunters Pt. S.E.</u>					Container type <u>2x6" Brass Tube</u>	ICED	Preservative	Sampling method <u>6" Hollow Stem Auger</u>	TPH-Extraction	BTEX/Total Fuel HCs	601 / 8010 Halocarbons	602 / 8020 Aromatics	604 / 8040 Phenols	606 / 8080 Pesticides/PCBs	624-8240 Purgeables			625-8270 BNAs & Pest (SV/s)	Dioxins	Metals	Wet Chemistry
Sampler (signature) <u>Arjun Chemburkar & Cheryl Seath</u>																					
Printed name <u>ARJUN CHEMBURKAR & CHERYL SEATH</u>																					
Lab Report Recipient																					
Telephone No.																					
Receiving Lab <u>CENTRAL COAST ANALYTICAL</u>																					
Address <u>SUBURBAN BD. STE. C-4</u>																					
<u>SAN LUIS OBISPO, CA. 93401</u>																					
Sample ID#	Time	W-water S-soil	C-comp G-grab	Volume																	
7672 3D-18 (1.5-2)	1038	S	G	1/2 lb.																	
7673 3C-19 (2-2.5)	1056																				
7674 3C-17 (2-2.5)	1113																				
7675 3B-18 (2-2.5)	1129																				
7676 3B-16 (2-2.5)	1142																				
7677 3B-14 (2-2.5)	1153																				
7678 3A-13 (2-2.5)	1205																				
7679 3A-11 (2-2.5)	1218																				
7680 3B-6 (2-2.5)	1228																				
7681 3A-9 (1.5-2)	1141																				

5 DAY
Turn
Around!

PCB ^{by} EPA
method 8080
and
% Moisture

Precautions: Conc: ☒ Lo ☐ Med ☐ HI Ship Via Bus Total Number of Containers: 10

Sample Relinquished By	Date	Time	Received By	Date	Time	Reason for Transfer (List Shipping Bill Number)
<u>Cheryl Seath</u> Company <u>ERM-West</u>	<u>8/9/88</u>	<u>0914</u>	<u>Lani Farlick</u> Company	<u>8/9/88</u>	<u>13:00</u>	
<u>Cheryl Seath (for Arjun)</u> Company <u>ERM-West</u>	<u>8/8/88</u>	<u>1900</u>				

LABORATORY: ☐ Samples Intact ☐ Samples at 4°C ☐ Samples not leaking ☐ # of containers matches C-of-C ☐ Container tags match C-of-C ☐ Cooler seals Intact

Please Complete Lab sample custodian Signature _____ Date _____ Time _____ Sample Disposition ☐ Return to Site ☐ Discard ☐ Hold _____ days

ERI-West

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Date 8/8/88 Weather cool, windy, sunny Page 2 of 2

Chain of Custody Record

Job # <u>40058</u>		Collection			GC		GC/MS		Inorg	Other	Number of Containers	Remarks									
Job Location <u>Hunters Pt. S.F.</u>		Container type	ICED	Preservative	Sampling method	TPH-Extraction	BTEX/Total Fuel HCs	601 / 8010 Halocarbons	602 / 8020 Aromatics	604 / 8040 Phenols			608 / 8080 Pest/PCBs	624-8240 Purgeables	625-8270 BNAs & Pest (SVs)	Dioxins	Metals	Wet Chemistry			
Sampler (signature) <u>Cheryl E. Seath & Arun Chemburkar</u>													<u>5 DAY</u> <u>TURN</u> <u>AROUND!</u>								
Printed name <u>Cheryl E. Seath & Arun Chemburkar</u>																					
Lab Report Recipient																					
Telephone No.																					
Receiving Lab <u>Central Coast Analytical</u>																					
Address <u>141 SUBURBAN RD. STE. C-4</u>													<u>TEST METHOD:</u> <u>EPA 8080</u> <u>FOR PCB'S.</u> <u>ANAL %</u> <u>MOISTURE</u>								
<u>SAN LUIS OBISPO, CA. 93401</u>																					
Sample ID#	Time	W-water S-sol	C-comp G-grab	Volume	Container type	ICED	Preservative	Sampling method	TPH-Extraction	BTEX/Total Fuel HCs	601 / 8010 Halocarbons	602 / 8020 Aromatics		604 / 8040 Phenols	608 / 8080 Pest/PCBs	624-8240 Purgeables	625-8270 BNAs & Pest (SVs)	Dioxins	Metals	Wet Chemistry	
7682 <u>3C-5</u> <u>(1.5-2')</u>	<u>1342</u>	<u>S</u>	<u>G</u>	<u>1.5 lb.</u>	<u>2x6" Brass Tubes</u>	<u>Y</u>	<u>N</u>	<u>CALIF. MODIFIED</u>							<u>X</u>						
7683 <u>3D-4</u> <u>(2.5-3')</u>	<u>1352</u>														<u>X</u>						
7684 <u>3E-5</u> <u>(2-2.5')</u>	<u>1406</u>														<u>X</u>						
7685 <u>3F-4</u> <u>(2-2.5')</u>	<u>1420</u>														<u>X</u>						
7686 <u>3G-5</u> <u>(2.5-3')</u>	<u>1435</u>														<u>X</u>						
7687 <u>3G-7</u> <u>(2.5-3')</u>	<u>1451</u>														<u>X</u>						
7688 <u>3H-6</u> <u>(2-2.5')</u>	<u>1506</u>														<u>X</u>						
7689 <u>3H-8</u> <u>(2-2.5')</u>	<u>1520</u>													<u>X</u>							
7690 <u>3J-9</u> <u>(2.5-3')</u>	<u>1608</u>													<u>X</u>							
7691 <u>3K-10</u> <u>(2.5-3')</u>	<u>1620</u>													<u>X</u>							
Precautions:		Conc: <input checked="" type="checkbox"/> Lo <input type="checkbox"/> Med <input type="checkbox"/> HI			Ship Via <u>Bus</u>		Total Number of Containers:		<u>10</u>												
Sample Relinquished By	Date	Time	Received By	Date	Time	Reason for Transfer (List Shipping Bill Number)															
<u>Cheryl E. Seath</u> Company <u>ERM-West</u>			<u>Gene Hartwick</u> Company	<u>8/8/88</u>	<u>1500</u>																
<u>Cheryl Seath (for Asc)</u> Company <u>ERM-West</u>	<u>8/8/88</u>	<u>81900</u>	Company																		
LABORATORY: <input type="checkbox"/> Samples Intact <input type="checkbox"/> Samples at 4°C <input type="checkbox"/> Samples not leaking <input type="checkbox"/> # of containers matches C-of-C <input type="checkbox"/> Container tags match C-of-C <input type="checkbox"/> Cooler seals intact																					
Please Complete Lab sample custodian Signature _____ Date _____ Time _____ Sample Disposition <input type="checkbox"/> Return to Site <input type="checkbox"/> Discard <input type="checkbox"/> Hold _____ days																					

7-20-88 D.J.

ER-I-West

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Date 8/8/88 Weather cool, windy, sunny Page 3 of 3

Chain of Custody Record

E

7692

7693

7694

7695

7696

7697

7698

7699

Job # <u>40058</u>					Collection				GC				GC/MS		Inorg	Other	Remarks <u>5 DAY</u> <u>TURN</u> <u>AROUND</u> <u>TIME!</u>					
Job Location <u>Hunters Pt. D.F.</u>					Container type	ICED	Preservative	Sampling method	TPH-Extraction	BTEX/Total Fuel HCs	601 / 8010 Halocarbons	602 / 8020 Aromatics	604 / 8040 Phenols	606 / 8080 Pest/PCBs	624-8240 Purgeables	625-8270 BNAs & Pest (SVs)		Dioxins	Metals	Wet Chemistry		
Sampler (signature) <u>Arun Chemburkar & Cheryl Seath</u>																						
Printed name <u>Arun Chemburkar & Cheryl Seath</u>																						
Lab Report Recipient																						
Telephone No.					Number of Containers																	
Receiving Lab <u>Central Coast Analytical</u>																						
Address <u>141 SUBURBAN RD. STE C4</u>																						
<u>SAN LUIS OBISPO, CA 93401</u>																						
Sample ID#	Time	W-water S-soil	Calcomp G-grab	Volume	Container type	ICED	Preservative	Sampling method	TPH-Extraction	BTEX/Total Fuel HCs	601 / 8010 Halocarbons	602 / 8020 Aromatics	604 / 8040 Phenols	606 / 8080 Pest/PCBs	624-8240 Purgeables	625-8270 BNAs & Pest (SVs)	Dioxins	Metals	Wet Chemistry	Number of Containers	EPA Protocol:	
3K-12 (2.5-3')	1645	S	G	~.5 lb.	3x6" Brass Tube	Y	N	6" Hollow Stem Auger						X							1	PCB Method 8080 and % Moisture
3F-16 (2.0-2.5')	1657	S	G	"	"	Y	N	"						X							1	
3E-17 (2-2.5')	1713	S	G	"	"	Y	N	"						X							1	
3G-15 (1.5-2')	1605	S	G	.25 lb.	1x6" Brass Tube	Y	N	Hand Auger						X							1	
3H-12 (1.5-2')	1625				"	Y	N	"						X							1	
3J-13 (1.5-2')	1640				"	Y	N	"						X							1	
3H-14 (1.5-2')	1702				"	Y	N	"						X							1	
3D-16 (1.5-2')	1715				"	Y	N	"						X							1	
Precautions:					Conc: <input checked="" type="checkbox"/> Lo <input type="checkbox"/> Med <input type="checkbox"/> HI					Ship Via					Total Number of Containers: <u>8</u>							
Sample Relinquished By		Date	Time	Received By		Date	Time	Reason for Transfer (List Shipping Bill Number)														
<u>Cheryl Seath</u> Company <u>ERH-West</u>		<u>8/8/88</u>	<u>0914</u>	<u>Kimi Hardick</u> Company		<u>8/8/88</u>	<u>1500</u>															
<u>Cheryl Seath (for Arun)</u> Company <u>ERH-West</u>		<u>8/8/88</u>	<u>1900</u>																			
LABORATORY—		<input type="checkbox"/> Samples Intact		<input type="checkbox"/> Samples at 4°C		<input type="checkbox"/> Samples not leaking		<input type="checkbox"/> # of containers matches C-of-C		<input type="checkbox"/> Container tags match C-of-C		<input type="checkbox"/> Cooler seals Intact										
Please Complete		Lab sample custodian		Signature		Date		Time		Sample Disposition		<input type="checkbox"/> Return to Site		<input type="checkbox"/> Discard		<input type="checkbox"/> Hold _____ days						

APPENDIX F

LABORATORY QUALITY CONTROL ASSURANCE
AND QUALITY CONTROL PROCEDURES

Central
Coast
Analytical
Services

**CENTRAL COAST
ANALYTICAL SERVICES**

Air, Water & Hazardous Waste Analysis
141 Suburban Road, Suite C-4
San Luis Obispo, California 93401
(805) 543-2553

RECEIVED
NOV 13 1987

ERM-WEST
WALNUT CREEK, CA

August 1, 1986

QUALITY ASSURANCE/QUALITY CONTROL PROCEDURE

1. Chain of Custody samples are stored in locked refrigerators.
2. The assignment of unique laboratory numbers to each sample as it is logged in has been extended to include the assignment of unique log numbers to subsamples on a container-by-container basis.
3. A central standard logbook is maintained for all standards. Information such as suppliers, lot numbers, weight/volume of standards used, date prepared and the name of the analyst preparing the standard are all part of the records kept therein.
4. The laboratory runs calibration standards at a minimum of three concentrations. We bracket sample data with standards.
5. We confirm all positive gas chromatographic results by either a second column or by GC/MS.
6. Blanks, duplicates, and spikes are analyzed once per batch, once per matrix type or once per 20 samples, whichever is more frequent. We have adopted an "all with the final report" filing system.
7. Records of the analysis of blank samples are recorded in individual laboratory notebooks, referred to on laboratory worksheets and documented on instrument use record books.
8. Acceptance limits for quality control samples currently used by our laboratory are summarized in Table I.
9. All analytical and quality control results are reviewed and approved by a supervisor. Data of an unusual nature are brought to the attention of one of our three Ph.D. chemists for final review.
10. Instructions for corrective action in the event of an out-of-control method are as follows: Stop analyses. Conduct investigation. Check mathematics. Check dilutions for systematic error. Check syringes, automatic pipettes, ect. for possible malfunction. Check dates on standards. Remake standards. If instrument malfunction is indicated, arrange for

service. Analyses may resume only when problem has been identified and corrected.

11. Repair and maintenance records are documented for inspection and review.
12. Analytical results including raw data and chromatographs of method blanks, three point standard calibration and quality control samples (matrix spikes and matrix spike duplicates) for the following methods are available for review:

<u>Method Number</u>	<u>Description</u>
632	Carbamates
8030	Acrolein, Acrylonitrile & Acetonitrile
8060	Phthalate Esters
8090	Nitroaromatics & Cyclic Ketones
8120	Chlorinated Hydrocarbons

TABLE I - ACCEPTANCE LIMITS for QUALITY ASSURANCE SAMPLES

ANALYTICAL PROCEDURE or PARAMETER GROUP		SPIKE LEVEL	ACCEPTABLE % RANGE	DUPLICATE ACCEPTABLE % DIFFERENCE
	Matrix	ppb		
Metals by AA *	Liquid	10X	80-115	15
	Solid		65-130	30
As	Liquid	468	70-120	25
	Solid	12500	60-135	35
Cd	Liquid	90	80-122	25
	Solid	12500	60-135	35
Cr	Liquid	916	75-120	20
	Solid	20000	60-135	35
Co	Liquid	303	82-115	15
	Solid	12500	60-135	40
Cu	Liquid	110	84-117	15
	Solid	180000	65-130	40
Pb	Liquid	556	85-120	15
	Solid	12000	75-125	25
Hg	Liquid	24	65-133	30
	Solid	6100	40-150	45
Ni	Liquid	121	82-118	15
	Solid	9000	55-140	40
Se	Liquid	45	50-131	40
	Solid	12500	40-150	45
V	Liquid	791	82-123	25
	Solid	9000	55-140	40
Zn	Liquid	1378	85-114	15
	Solid	360000	40-150	45
Tl	Solid	1800	50-145	40
Ag	Solid	1800	12-160	45
Sb	Solid	6100	60-135	35
Be	Solid	16300	55-140	40
Al	Liquid	815	78-125	25

TABLE I - ACCEPTANCE LIMITS for QUALITY ASSURANCE

SAMPLES-Continued

ANALYTICAL PROCEDURE or PARAMETER GROUP		SPIKE LEVEL ACCEPTABLE		DUPLICATE
Matrix	ppb	% RANGE	% DIFFERENCE	
Chlorinated				
Herbicides *	Liquid	300	75-120	25
2,4-D	Liquid	300	75-120	25
Silvex	Liquid	70	75-120	25
2,4,5-T	Liquid	90	75-125	25
Phenols *				
	Liquid	300	40-140	45
	Solid			
2-Chlorophenol	Liquid	100	25-135	45
	Solid	10030	32-140	50
2-Nitrophenol	Liquid	500	D-150	100
	Solid	4020	D-150	100
Phenol	Liquid	300	5-112	75
	Solid	4000	D-150	100
2,4-Dimethyl-phenol	Liquid	300	12-150	75
	Solid	4020	D-150	100
2,4-Dichloro-phenol	Liquid	300	35-140	50
	Solid	10100	19-150	75
2,4,6-Tri-chlorophenol	Liquid	300	20-150	75
	Solid	10020	20-150	75
4-Chloro-3-methylphenol	Liquid	200	22-147	75
	Solid	10020	15-150	75
2,4-Dinitro-phenol	Liquid	500	D-160	100
	Solid	10000	D-150	100
2-Methyl-4,6-dinitrophenol	Liquid	500	5-160	75
	Solid	4020	D-160	100
Pentachloro-phenol	Liquid	100	14-176	75
	Solid	4010	D-180	100
4-Nitrophenol	Liquid	500	D-132	75
	Solid	4010	D-160	100

- TABLE I - ACCEPTANCE LIMITS for QUALITY ASSURANCE SAMPLES

- Continued

ANALYTICAL PROCEDURE or PARAMETER GROUP		SPIKE LEVEL ACCEPTABLE		DUPLICATE
	Matrix	ppb	% RANGE	ACCEPTABLE % DIFFERENCE

Metals by ICP *	Liquid	10X	50-140	40
As	Liquid	22	70-125	25
Cd	Liquid	2.5	52-140	40
Cr	Liquid	10	46-145	40
Cu	Liquid	11	10-160	75
Pb	Liquid	24	15-155	75
Ni	Liquid	60	58-135	40
Se	Liquid	6	D-200	100
Zn	Liquid	16	10-170	75
Be	Liquid	20	70-125	25
Co	Liquid	120	40-150	50
V	Liquid	70	80-115	15
Carbamate by LC	Liquid	100X	80-115	15
Polychlorinated Biphenyls (see organochlorine pesticides)				
Aromatic Volatile Organics				
Benzene	Liquid	9	64-130	40
Chlorobenzene	Liquid	100	70-127	30
1,2-Dichloro- benzene	Liquid	10	16-170	75
1,3-Dichloro- benzene	Liquid	5	37-150	75
1,4-Dichloro- benzene	Liquid	10	40-170	75
Ethylbenzene	Liquid	10	62-137	40
Toluene	Liquid	100	41-136	75

TABLE I - ACCEPTANCE LIMITS for QUALITY ASSURANCE SAMPLES
- Continued

ANALYTICAL PROCEDURE or PARAMETER GROUP		SPIKE LEVEL ACCEPTABLE		DUPLICATE ACCEPTABLE
Matrix		ppb	% RANGE	% DIFFERENCE

Halogenated Volatile Organics				
Bromodichloro-				
methane	Liquid	0.5	85-115	15
Bromoform	Liquid	50	60-130	30
Carbon				
tetrachloride	Liquid	50	20-140	75
Chlorobenzene	Liquid	50	67-121	25
Chloroethane	Liquid	50	30-150	75
2-Chloroethyl				
vinyl ether	Liquid	133	66-126	30
Chloroform	Liquid	50	60-160	40
Chloromethane	Liquid	24	50-140	50
Dibromochloro-				
methane	Liquid	93	78-118	25
1,2-Dichloro-				
benzene	Liquid	150	85-115	15
1,3-Dichloro-				
benzene	Liquid	50	79-112	25
1,4-Dichloro-				
benzene	Liquid	50	70-124	25
1,1-Dichloro-				
ethane	Liquid	45	80-120	25
1,2-Dichloro-				
ethane	Liquid	45	80-120	25
1,1-Dichloro-				
ethylene	Liquid	50	40-150	60
t-1,2-Dichloro-				
ethylene	Liquid	98	30-150	75
1,2-Dichloro-				
propane	Liquid	39	70-125	25
t-1,3-Dichloro-				
propylene	Liquid	50	22-150	75
1,1,2,2-Tetra-				
chloroethane	Liquid	45	45-135	45
Tetrachloro-				
ethylene	Liquid	35	35-150	75
1,1,1-Tri-				
chloroethane	Liquid	29	45-135	50
1,1,2-Tri-				
chloroethane	Liquid	50	20-150	75
Trichloro-				
ethylene	Liquid	45	80-120	25
Vinyl Chloride	Liquid	32	70-130	25

TABLE I - ACCEPTANCE LIMITS for QUALITY ASSURANCE SAMPLES
- Continued

ANALYTICAL PROCEDURE or PARAMETER GROUP		SPIKE		DUPLICATE
Matrix		LEVEL	ACCEPTABLE	ACCEPTABLE
		ppb	% RANGE	% DIFFERENCE

Base/Neutrals				
Acenapthene	Liquid	100	47-145	45
	Solid	2010	26-130	75
Acenapthalene	Liquid	2000	31-140	75
	Solid	2020	25-130	75
Anthracene	Liquid	2000	30-140	75
	Solid	4040	32-132	75
Benzo(a)anth- racene	Liquid	2000	30-140	75
	Solid	2020	7-150	100
Benzo(b)fluor- anthene	Liquid	2000	17-140	100
	Solid	4420	6-150	100
Benzo(k)fluor- anthene	Liquid	2000	17-140	100
	Solid	4420	6-150	100
Benzo(ghi)per- ylene	Liquid	2000	D-180	200
	Solid	62000	40-160	75
Benzo(a)pyrene	Liquid	2000	17-140	100
	Solid	2080	18-130	100
Benzidine	Liquid	2000	0-150	200
	Solid	2010	EPA NOT ABLE TO RECOVER	
Butyl benzyl phthalate	Liquid	2000	D-150	200
	Solid	2020	7-150	100
Bis(2-chloro- ethoxy)methane	Liquid	2000	D-150	200
	Solid	2020	9-150	100
Bis(2-chloro- ethyl)ether	Liquid	2000	D-150	200
	Solid	2020	D-150	200
Bis(2-chloro- ispropyl)ether	Liquid	2000	D-150	200
	Solid	10020	30-150	75
Bis(2-ethyl- hexyl)phthal.	Liquid	2000	D-200	200
	Solid	32210	30-150	75
4-Bromophenyl phenyl ether	Liquid	2000	30-150	75
	Solid	2070	30-150	75
2-Chloro- naphthalene	Liquid	2000	37-150	75
	Solid	1920	30-150	75
4-chlorophenyl phenyl ether	Liquid	2000	30-150	75
	Solid	2010	15-130	100
Chrysene	Liquid	2000	37-150	75
	Solid	2020	7-150	100
Dibenzo(a,h) anthracene	Liquid	2000	D-170	200
	Solid	61400	34-200	100
Di-n-butyl phthalate	Liquid	2000	D-200	200
	Solid	2010	5-180	100
1,2-Dichloro benzene	Liquid	100	1-118	125
	Solid	77200	50-128	50

TABLE I - ACCEPTANCE LIMITS for QUALITY ASSURANCE SAMPLES
- Continued

ANALYTICAL PROCEDURE or PARAMETER GROUP		SPIKE LEVEL	ACCEPTABLE % RANGE	DUPLICATE ACCEPTABLE % DIFFERENCE
	Matrix	ppb		
1,3-Dichloro	Liquid	100	1-118	125
benzene	Solid	8540	D-170	200
1,3-Dichloro	Liquid	100	1-118	125
benzene	Solid	54800	29-142	75
3,3-Dichloro-	Liquid	2000	D-300	200
benzidine	Solid	10000	D-300	200
Dimethyl	Liquid	2000	3-130	125
phthalate	Solid	2010	21-140	100
2,4-Dinitro-	Liquid	200	50-158	75
toluene	Solid	2030	18-140	100
2,6-Dinitro-	Liquid	200	50-158	75
toluene	Solid	2030	18-140	100
Di-n-octyl	Liquid	2000	D-200	200
phthalate	Solid	2030	12-170	100
Fluoranthene	Liquid	2000	20-140	100
	Solid	2010	D-170	150
Fluorene	Liquid	2000	20-140	100
	Solid	2010	30-160	100
Hexachloro-	Liquid	2000	26-150	100
benzene	Solid	1610	63-133	50
Hexachloro-	Liquid	2000	D-200	200
butadiene	Solid	2050	0-200	200
Hexachloro-	Liquid	2000	D-200	200
cypentadiene	Solid	2010	EPA NOT ABLE TO RECOVER	
Hexachloro-	Liquid	2000	D-200	200
ethane	Solid	2000	56-151	75
Indeno(1,2,3-	Liquid	2000	D-200	200
cd) pyrene	Solid	50010	34-180	100
Isophorone	Liquid	2000	D-200	200
	Solid	2020	9-142	100
Naphthalene	Liquid	2000	D-200	200
	Solid	152500	55-126	75
Nitrobenzene	Liquid	2000	D-200	200
	Solid	42100	16-150	125
N-Nitrosodi-n-	Liquid	2000	D-200	200
propylamine	Solid	10020	62-180	75
N-Nitrosodi-	Liquid	2000	6-170	100
phenylamine	Solid	10000	0-200	200
Phenanthrene	Liquid	2000	29-136	100
	Solid	2010	26-130	75
Pyrene	Liquid	2000	29-136	100
	Solid	2020	16-160	100
Trichloro-	Liquid	2000	17-170	100
benzene	Solid	2200	3-160	125

TABLE I - ACCEPTANCE LIMITS for QUALITY ASSURANCE SAMPLES
- Continued

ANALYTICAL PROCEDURE or PARAMETER GROUP		SPIKE LEVEL ACCEPTABLE		DUPLICATE ACCEPTABLE
Matrix		ppb	% RANGE	% DIFFERENCE
EPA 624/8240 Purgeable Organics				
Benzene	Liquid	100	68-130	40
	Solid	99	80-100	50
Bromodichloro-	Liquid	100	73-130	30
methane	Solid	10	60-170	50
Bromoform	Liquid	100	52-152	50
	Solid	10	70-150	50
Bromomethane	Liquid	100	22-169	75
Carbon tetra-	Liquid	100	69-149	40
chloride	Solid	10	61-150	50
Chlorobenzene	Liquid	100	75-130	30
	Solid	10	10-150	100
Chloroethane	Liquid	100	14-190	100
2-Chloroethyl				
vinyl ether	Liquid	100	75-150	35
Chloroform	Liquid	100	64-136	40
	Solid	10	85-132	40
Chloromethane	Liquid	100	28-172	75
Dibromochloro-	Liquid	100	62-146	40
methane	Solid	1	33-126	75
1,1-Dichloro-	Liquid	100	60-150	40
ethane	Solid	10	31-150	75
1,2-Dichloro-	Liquid	100	72-132	35
ethane	Solid	10	13-160	100
1,1-Dichloro-	Liquid	100	54-145	40
ethene	Solid	10	10-170	100
1,2-Dichloro-	Liquid	100	71-130	35
ethene	Solid	10	23-147	100
1,2-Dichloro-	Liquid	100	67-136	40
propane	Solid	10	35-144	75
1,3-Dichloro-	Liquid	100	45-157	50
propene	Solid	8	45-140	75
1,3-Dichloro-	Liquid	100	46-154	50
propene	Solid	10	48-142	100
Ethylbenzene	Liquid	100	73-133	40
	Solid	10	43-130	100
Methylene	Liquid	100	8-187	200
Chloride	Solid	10	28-130	200
1,1,2,2-Tetra-	Liquid	100	60-146	40
chloroethane	Solid	10	D-250	300
Tetrachloro-	Liquid	100	67-133	70
ethene	Solid	10	D-250	300

TABLE I - ACCEPTANCE LIMITS for QUALITY ASSURANCE SAMPLES
- Continued

ANALYTICAL PROCEDURE or PARAMETER GROUP		SPIKE LEVEL ACCEPTABLE		DUPLICATE ACCEPTABLE
Matrix		ppb	% RANGE	% DIFFERENCE
Toluene	Liquid	100	56-140	40
	Solid	10	61-120	45
1,1,1-Tri-	Liquid	100	54-150	40
chloroethane	Solid	10	23-140	150
1,1,2-Tri-	Liquid	100	59-149	40
chloroethane	Solid	10	55-127	50
Trichloro-	Liquid	100	64-136	40
ethene	Solid	10	18-145	150
Trichloro-	Liquid	100	50-164	75
fluoromethane				
Vinyl chloride	Liquid	100	23-173	180
EPA 8030 PURGEABLE ORGANIC COMPOUNDS				
Acrolein	Liquid	20	61-135	45
Acrylonitrile	Liquid	20	85-124	40
EPA 8060 PHTHALATE ESTERS				
Bis(2-ethyl-	Liquid	1000	74-111	45
hexyl)phthalate				
Butyl benzyl	Liquid	1000	66-116	50
phthalate				
Dibutyl-	Liquid	1000	65-115	50
phthalate				
Diethyl	Liquid	50	90-110	40
phthalate				
Dimethyl	Liquid	50	84-110	40
phthalate				
Dioctyl	Liquid	150	73-113	40
phthalate				
EPA 8090 NITROAROMATICS AND CYCLIC KETONES				
2,4-Dinitro-	Liquid	100	55-110	75
toluene				
2,6-Dinitro-	Liquid	50	60-110	75
Isophorone	Liquid	50	63-110	75
toluene				
Nitrobenzene	Liquid	100	58-113	75
EPA 8100 POLYNUCLEAR AROMATIC HYDROCARBONS				
Same as EPA 625/8270				

TABLE I - ACCEPTANCE LIMITS for QUALITY ASSURANCE SAMPLES
- Continued

ANALYTICAL PROCEDURE or PARAMETER GROUP		SPIKE LEVEL ACCEPTABLE		DUPLICATE ACCEPTABLE
Matrix		ppb	% RANGE	% DIFFERENCE
EPA 8120 CHLORINATED HYDROCARBONS				
2-Chloro-naphthalene	Liquid	200	19-157	180
1,2-Dichloro-benzene	Liquid	300	57-125	100
1,3-Dichloro-benzene	Liquid	200	40-146	150
1,4-Dichloro-benzene	Liquid	300	35-153	150
Hexachloro-benzene	Liquid	10	61-134	75
Hexachloro-butadiene	Liquid	30	66-130	75
Hexachloro-ethane	Liquid	10	63-136	75
1,2,4-Tri-chlorobenzene	Liquid	10	48-148	125
EPA 8140 ORGANOPHOSPORUS PESTICIDES				
Azinphos methyl	Liquid	200	32-141	150
Bolstar	Liquid	40	53-115	75
Chlorpyrifos	Liquid	50	82-116	45
Coumaphos	Liquid	200	70-147	75
Demeton	Liquid	300	46-121	100
Diazinon	Liquid	5	55-112	75
Dichlorvos	Liquid	500	55-117	75
Disulfoton	Liquid	80	60-122	75
Ethoprop	Liquid	50	88-112	45
Fensulfothion	Liquid	100	46-148	150
Fenthion	Liquid	50	33-136	150
Merphos	Liquid	50	76-145	75
Mevinphos	Liquid	500	34-123	150
Naled	Liquid	200	69-119	75
Parathion methyl	Liquid	500	81-115	45
Phorate	Liquid	40	46-117	150
Ronnel	Liquid	50	83-117	45
Stiophos	Liquid	500	54-112	150
Tokuthion	Liquid	50	51-113	150
Trichloronate	Liquid	20	46-161	150

90, 96-98, and 104-115 to new Table IB, entitled "List of Approved Inorganic Test Procedures", adding two new organic parameters, Carbonaceous Biochemical Oxygen Demand (CBOD₅) and Nitrate-Nitrite, including an additional test procedure based upon the inductively coupled plasma technique in Table IB for 25 of the metal parameter designations, by including 10 methods approved under the equivalency provisions of §§ 136.4(d) and 136.5(e), and updating references to EPA, Standard Methods, ASTM, AOAC and USGS test procedures; by deleting

former parameter 14 (Chlorinated organic compounds) and by entering the individual chlorinated organic compounds into new Table IC, entitled, "List of Approved Test Procedures for Non-Pesticide Organic Compounds", transferring old parameters 9 (Benzidine) and 94 (Pentachlorophenol) to Table IC, by including the 78 additional proposed non-pesticidal organic parameters and by adding 17 new test procedures in Table IC; by deleting former parameter 95 (Pesticides) and by entering the 68 individual pesticides into new Table ID,

entitled "List of Approved Test Procedures for Pesticides", by including the 2 additional proposed pesticide parameters, and the two new test procedures in Table ID; and by transferring the former radiological parameters 100-103 to new Table IF, entitled "Approved Radiological Test Procedures", adding an EPA reference to the approved test procedures, and updating the Standard Methods, ASTM and USGS references. As revised, Table I reads as follows:

§ 136.3 Identification of Test Procedures.

TABLE IA.—LIST OF APPROVED BIOLOGICAL TEST PROCEDURES

Parameter and units	Method ¹	EPA ²	Reference (Method Number or Page)		
			Standard Methods 15th Ed.	ASTM	USGS
Section:					
1. Coliform (fecal) number per 100 ml	MPN, 5 tube, 3 dilution, or, membrane filter (MF) ³ , single step	p. 132 p. 124	980C 908C		B-0050-77.
2. Coliform (fecal) in presence of chlorine number per 100 ml	MPN, 5 tube, 3 dilution	p. 132	908C		
3. Coliform (total) number per 100 ml	MPN, 5 tube, 3 dilution, or, MF ³ single step or two step	p. 114 p. 108	908A 909A		B-0025-77.
4. Coliform (total) in presence of chlorine, number per 100 ml	MPN, 5 tube, dilution, or MF ³ with enrichment	p. 114 p. 111	908A 909 (A + A.5c)		
5. Fecal streptococci, number per 100 ml	MPN, 5 tube, 3 dilution; MF ³ ; or, plate count	p. 139 p. 136 p. 143	910A 910B 910C	B0055-77. ⁴	

Table IA Notes

- ¹ The method must be specified when results are reported.
- ² "Microbiological Methods for Monitoring the Environment, Water and Wastes, 1978", EPA-600/8-78-017, U.S. Environmental Protection Agency.
- ³ Green, P.E., et al., Methods for Collection and Analysis of Aquatic Biological and Microbiological Samples, "U.S. Geological Survey, Techniques of Water-Resources Investigations, Book 5, Chapter A4, Laboratory Analysis, 1977
- ⁴ D 45 um membrane filter or other pore size certified by the manufacturer to fully retain organisms to be cultivated, and free of extractables which could interfere with their growth and development.
- ⁵ Approved only if dissolution of the KF Streptococcus Agar (Section 5.1, USGS Method B-0055-77) is made in a boiling water bath to avoid scorching of the medium.

TABLE IB.—LIST OF APPROVED INORGANIC TEST PROCEDURES

Parameter, units, and method	Reference (method No. or page)				
	EPA 1978	Standard methods 15th Ed.	ASTM	USGS ¹	Other
1. Acidity, as CaCO ₃ , mg/L. Electrometric end point or phenolphthalein end point	305.1	402(4 d)	D1067-70(E)		
2. Alkalinity, as CaCO ₃ , mg/L. Electrometric or colorimetric					
Titration to pH 4.5, manual	310.1	403	D1067(B)	I-1030-78	P. 548. ²
Or automated	310.2			I-2030-78	
3. Aluminum—Total ³ , mg/L. Digestion ³ followed by AA direct aspiration	202.1	303C		I-3051-78	Method 200.7. ⁴
AA furnace	202.2	304			
Inductively coupled plasma					
Or colorimetric (Eriochrome cyanine R)		306B			
4. Ammonia (as N), mg/L. Manual distillation ⁵ (at pH 9.5):					
Followed by	350.2	417A			
Nesslerization	350.2	417B	D1426-79(A)	I-3520-78	P. 553. ²
Distillation	350.2	417D			
Electrode	350.3		D1426-79(D)		
Automated phenate or automated electrode	350.1	417F	D1426-79(C)	I-4523-78	
5. Arsenic—Total ³ , mg/L. Digestion ³ followed by AA direct aspiration	204.1	303A			Method 200.7. ⁴
AA furnace, or	204.2	304			
Inductively coupled plasma					
6. Arsenic—Total ³ , mg/L. Digestion ³ followed by hydride	206.5				
AA furnace	206.3	303E	D2972-79(B)	I-3062-78	Method 200.7. ⁴
Inductively coupled plasma	206.2	304			
Or, colorimetric (SODC)	206.4	307B	D2972-79(A)	I-3080-78	
7. Barium—Total ³ , mg/L. Digestion ³ followed by AA direct aspiration	208.1	303C		I-3084-78	Method 200.7. ⁴
AA furnace, or	208.2	304			
Inductively coupled plasma					
8. Beryllium—Total ³ , mg/L. Digestion ³ followed by AA direct aspiration	210.1	303C	D3645-78	I-3085-78	
AA furnace	210.2	304			

TABLE IB.—LIST OF APPROVED INORGANIC TEST PROCEDURES—Continued

Parameter, units, and method	Reference (method No. or page)				
	EPA 1978	Standard methods 15th Ed.	ASTM	USGS ¹	Other
Inductively coupled plasma Or colorimetric (aluminum)		309B			Method 200.7. ^a
9. Biochemical oxygen demand (BOD ₅), mg/L Winkler (Azide modification) Or electrode method	405.1	507		I-1578-78	P. 17. ^a P. 548. ^a
10. Boron—Total, mg/L: Colorimetric (curcumin) or Inductively coupled plasma	212.3	404A		I-3112-78	Method 200.7. ^a P. 844. ¹⁰
11. Bromide, mg/L: Titrimetric	320.1		D1246-77(C)	I-1125-78	
12. Cadmium—Total ^a , mg/L: Digestion ^a followed by: AA direct aspiration AA furnace Inductively coupled plasma Volametry ^a or Colorimetric (dithione)	213.2 213.2	303A or 303B 304	D3557-78 (A or B)	I-3135-78 or I-3136-78	Pg. 557. ^a P. 37. ^a Method 200.7. ^a
13. Calcium—Total ^a , mg/L: Digestion ^a followed by Atomic absorption Inductively coupled plasma Or EDTA titration	215.1 215.2	303A 311C	D511-77(C) D511-77(B)	D152-78	Method 200.7. ^a
14. Carbonaceous Biochemical oxygen demand (CBOD ₅), mg/L: Winkler (Azide modification) or electrode method with nitrification inhibitor.		507(5 & 6)			
15. Chemical oxygen demand (COD), mg/L: Titrimetric colorimetric Manual or Automated Spectrophotometric	410.1 410.2 410.3 410.4	506A	D1252-78	I-3560-78 I-3562-78 I-3561-78	P. 550. ^a and P. 17. ^a and (11) (11)
16. Chloride, mg/L: Titrimetric (silver nitrate) or Mercuric nitrate Colorimetric (mercuric nitrate) manual or Automated	325.3 325.1 or 325.2	407A 407B 407D	D512-67(B) D512-67(A) D512-67(C)	I-1163-78 I-1164-78 I-1167-78 I-2187-78	P. 554. ^a
17. Chlorine—Total residual, mg/L: Titrimetric-spectrometric ¹¹ Starch end point Iodometric or DPD-FAS Spectrophotometric, DPD, or Electrode	330.1 330.2 330.3 330.4 330.5	408C 408B 408A 408D 408E	D1253-76(A) D1253-76(B)		(11)
18. Chromium VI dissolved, mg/L: 0.45 micron filtration with: Extraction and atomic absorption, or Colorimetric (Diphenylcarbazide)	218.4	303B		I-1232-78 I-1230-78	
19. Chromium—Total ^a , mg/L: Digestion ^a (optional extraction) followed by: AA direct aspiration AA furnace Inductively coupled plasma Or colorimetric (Diphenylcarbazide)	218.3 218.1 218.2	303A or 303B 304	D1687-77(D)	I-3236-78	P. 557. ^a Method 200.7. ^a
20. Cobalt—Total ^a , mg/L: Digestion ^a followed by: AA direct aspiration AA furnace, or Inductively coupled plasma	219.1 219.2	303A or 303B 304	D3558-77 (A or B)	I-3240-78 or I-3238-78	P. 37. ^a Method 200.7. ^a
21. Color, platinum Cobalt units or dominant wavelength hue, luminance, purity Colorimetric, ADMI Platinum cobalt, or Spectrophotometric	110.1 110.2 110.3	204D 204A 204B		I-1250-78	(11)
22. Copper—Total ^a , mg/L: Digestion ^a followed by: AA direct aspiration AA furnace Inductively coupled plasma Colorimetric (Neocupron) Bismuthimetric	220.1 220.2	303A or 303B 304	D1688-77 (D or E)	I-3271-78 or I-3270-78	P. 557. ^a and P. 37. ^a Method 200.7. ^a
23. Cyanide—Total mg/L: Manual distillation with MgCl ₂ Followed by titrimetric Manual or Automated ¹² spectrophotometric	335.2 335.2 335.2 335.3	412D 412B 412C 412D	D2036-75(A) D2036-75(A) D2036-75(B)	I-3300-78	P. 22. ^a
24. Cyanide amenable to chlorination, mg/L: Manual distillation with MgCl ₂ , followed by titrimetric, manual or automated ¹² spectrophotometric	335.1	412F			
25. Fluoride—Total, mg/L: Manual distillation ¹ Followed by manual or Automated electrode SPADNS Or automated complexone	340.2 340.1 340.3	413A 413B 413C 413E	U1179-72(B) D1179-72(A)	I-4327-78	
26. Gold—Total ^a , mg/L: Digestion ^a followed by: AA direct aspiration Or AA furnace	231.1 231.2	303A 304			
27. Hardness—Total as CaCO ₃ , mg/L: Automated colorimetric EDTA titration Inductively coupled plasma Or atomic absorption (sum of Ca and Mg as their respective carbonates)	130.1 130.2 215.1 242.1	314B 303A	D1126-67(B)	I-1338-78 I-3153-78 I-3448-78	Method 200.7. ^a

TABLE IB.—LIST OF APPROVED INORGANIC TEST PROCEDURES—Continued

Parameter, units, and method	Reference (method No. or page)				
	EPA 1979	Standard methods 15th Ed.	ASTM	USGS ¹	Other
28. Hydrogen ion (pH), pH units: Electrometric	150.1	423	D1293-78(A) or D1293-78(B)	I-1586-78	(^m)
Measurements; or automated electrode					
29. Iodine—Total ² , mg/L: Digestion ³ followed by: AA direct aspiration	235.1	303A			
Or AA furnace	235.2	304			
30. Iron—Total ² , mg/L: Digestion ³ followed by: AA direct aspiration	236.1	303A or 303B	D1068-77 (C or D)	I-3381-78	P. 557. ⁴
AA furnace	236.2	304			Method 200.7. ⁴ (^m)
Inductively coupled plasma					
Or colorimetric (Phenanthroline)		315B	D1068-77(A)		
31. Kjeldahl nitrogen—Total (as N), mg/L: Digestion and distillation	351.2	420A or B			P. 552. ⁴
Followed by titration	351.3	417D	D3580-77		
Reassimilation or	351.3	417B			
Electrode	351.3	417E		I-4551-78 I-4552-78	
Automated phenate	351.1				
Semi-automated block digester	351.2				
Or potentiometric	351.4				
32. Lead—Total ² , mg/L: Digestion ³ followed by: AA direct aspiration	239.1	303A or 303B	D3558-78 (A or B)	I-3388-78	P. 557. ⁴
AA furnace	239.2	304			Method 200.7. ⁴
Inductively coupled plasma					
Voltammetry ² or			D3558-78(C)		
Colorimetric (Dithione)		318B			
33. Magnesium—Total ² , mg/L: Digestion ³ followed by: Atomic absorption	242.1	303A	D511-77(B)	I-3447-78	P. 557. ⁴ Method 200.7. ⁴
Inductively coupled plasma					
Or gravimetric		318B	D511-77(A)		
34. Manganese—Total ² , mg/L: Digestion ³ followed by: AA direct aspiration	243.1	303A or 303B	D658-77 (B or C)	I-3454-78	P. 557. ⁴
AA furnace	243.2	304			Method 200.7. ⁴
Inductively coupled plasma					P. 564. ⁴ 18.
Or colorimetric (Persulfate)		319B	D658-77(A)		
Peroxide					
35. Mercury—Total ² , mg/L: Cold vapor, manual or	245.1	303F	D3223-78	I-3462-78	P. 558. ⁴
Automated	245.2				
36. Molybdenum—Total ² , mg/L: Digestion ³ followed by: AA direct aspiration	246.1	303C		I-3480-78	Method 200.7. ⁴
AA furnace, or	246.2	304			
Inductively coupled plasma					
37. Nickel—Total ² , mg/L: Digestion ³ followed by: AA direct aspiration	249.1	303A or 303B	D1886-77 (C or D)	I-3488-78	Method 200.7. ⁴
AA furnace	249.2	304			
Inductively coupled plasma					
Or colorimetric (Methylamine)		321B			
38. Nitrate (as N), mg/L: Bismite sulfate, or	352.1		D082-71		P. 554. ⁴
Nitrate-nitrite N minus Nitrite N	See parameters 39 and 40.	See parameters 39 and 40.	See parameters 39 and 40.	See parameters 39 and 40.	P. 28. ⁴
39. Nitrate-nitrite (as N), mg/L: Cadmium reduction, manual	352.3	418C	D3867-78(B)		
Or automated, or	352.2	418F	D3867-78(A)	I-4545-78	
Automated hydrazine	352.1				
40. Nitrite (as N), mg/L: Spectrophotometric, manual or	354.1	419	D1254-67		18.
Automated (Diazotization)				I-4540-78	
41. Oil and grease—Total recoverable, mg/L: Gravimetric (extraction)	413.1	503A			
42. Organic carbon—Total (TOC), mg/L: Combustion or oxidation	415.1	505	D2578-78(A) or D2578-78(B)		P. 551. ⁴ and P. 4. ⁴
43. Organic nitrogen (as N), mg/L: Total Kjeldahl N minus ammonia N	See parameters 31 and 4.	420A	D3580-77 minus D1428-78(A)	See parameters 31 and 4.	PP. 552-553. ⁴
44. Orthophosphate (as P), mg/L: Ascorbic acid method, automated	365.1	424G		I-4801-78	P. 561. ⁴
Or manual single reagent or	365.2	424F	D515-78(A)		
Manual two reagent	365.3				
45. Osmium—Total ² , mg/L: Digestion ³ followed by: AA direct aspiration, or	252.1	303C			
AA furnace	252.2	304			
46. Oxygen, dissolved, mg/L: Winkler (Azide modification)	360.2	421B	D1588-80(A)	I-1575-78	P. 550. ⁴
Or electrode	360.1	421F		I-1576-78	
47. Palladium—Total ² , mg/L: Digestion ³ followed by: AA direct aspiration	253.1				P. 527. ⁴
Or AA furnace	253.2				P. 528. ⁴
48. Phenols, mg/L: Manual distillation	420.1		D1783-70 (A or B)		26.
Followed by manual	420.1				26.
Or automated ¹¹ colorimetric (4AAP)	420.2				21.
49. Phosphorus (elemental), mg/L: Gas-liquid chromatography					

TABLE IB.—LIST OF APPROVED INORGANIC TEST PROCEDURES—Continued

Parameter, units, and method	Reference (method No. or page)				
	EPA 1979	Standard methods 15th Ed.	ASTM	USGS ¹	Other
50. Phosphorus—Total, mg/L Persulfate digestion Followed by manual or Automated ascorbic acid Reduction, or semi-automated block digester	365.2 365.2 or 365.3 365.1 365.4	424C (M) 424F 424G	D515-78(A)	I-4800-78 I-4803-78	P. 561. ²
51. Platinum—Total ³ , mg/L. Digestion ³ followed by: AA direct aspiration Or AA furnace	255.1 255.2	303A 304			
52. Potassium—Total ³ , mg/L. Digestion ³ followed by: Atomic absorption Inductively coupled plasma Or flame photometric	258.1	303A		I-3630-78	P. 560. ² Method 200.7. ⁴
53. Residue—total, mg/L. Gravimetric, 103-105°C	180.3	208A		I-3750-78	
54. Residue—filterable, mg/L. Gravimetric, 180°C	180.1	208B		I-1750-78	
56. Residue—nonfilterable, (TSS), mg/L. Gravimetric, 183-105°C post washing of residue	180.2	208D		I-3785-78	
58. Residue—settleable, mg/L. Volumetric (Imhoff cone) or gravimetric	180.5	208F			
57. Residue—volatile, mg/L. Gravimetric, 550°C	180.4	208E		I-3753-78	
59. Rhodium—Total ³ , mg/L. Digestion ³ followed by: AA direct aspiration Or AA furnace	265.1 267.2	303A 304			
60. Ruthenium—Total ³ , mg/L. Digestion ³ followed by: AA direct aspiration Or AA furnace	267.1 267.2	303A 304			
61. Selenium—Total ³ mg/L. Digestion ³ followed by: AA furnace Inductively coupled plasma Or hydride	270.2 270.3	304			Method 200.7. ⁴
62. Silver—Total ³ mg/L. Digestion ³ followed by: AA direct aspiration AA furnace, or Inductively coupled plasma	272.1 272.1	303A or 303B 304		I-3720-78	P. 557. ² and p. 37. ² Method 200.7. ⁴
63. Sodium—Total ³ , mg/L. Digestion ³ followed by: Atomic absorption Inductively coupled plasma Or flame photometric	273.1	303A		I-3735-78	P. 561. ² Method 200.7. ⁴
64. Specific conductance, mhos/cm. Wheatstone bridge	120.1	205	D1428-64(A) D1125-77(A)	I-1780-78	P. 547. ²
65. Sulfate (as SO ₄), mg/L: Automated methylthymol blue Gravimetric, or Turbidimetric	375.2 375.3 375.4	426A or 426B 426C	D516-68(A) D516-68(B)	I-2822-78	PP. 562-63. ²
66. Sulfide (as S), mg/L: Thermometric (iodine) or Colorimetric (methylene blue)	376.1 376.2	427D 427C		I-3840-78	
67. Sulfite (as SO ₃), mg/L. Thermometric (iodine iodate)	377.1	428F	D1339-78(C)		
68. Sulfite, mg/L. Colorimetric (methylene blue)	425.1	512A	D2330-68(A)		(79)
69. Temperature, °C. Thermometric	170.1	212			
70. Thallium—Total ³ mg/L. Digestion ³ followed by: AA direct aspiration AA furnace, or Inductively coupled plasma	279.1 279.2	303A 304			Method 200.7. ⁴
71. Tin—Total ³ mg/L. Digestion ³ followed by: AA direct aspiration or AA furnace	282.1 282.2	303A 304		I-3850-78	
72. Titanium—Total ³ mg/L. Digestion ³ followed by: AA direct aspiration or AA furnace	283.1 283.2	303C 304			
73. Turbidity, NTU. Nephelometric	180.1	214A	D1889-71	I-3980-78	
74. Vanadium—Total ³ mg/L. Digestion ³ followed by: AA direct aspiration AA furnace Inductively coupled plasma Or colorimetric (Gallate acid)	286.1 286.2	303C 304			Method 200.7. ⁴
75. Zinc—Total ³ mg/L. Digestion ³ followed by: AA direct aspiration AA furnace Inductively coupled plasma Or colorimetric (Zincon)	289.1 289.2	303A or 303B 304	D1691-77(D) D1691-77(C)	I-3900-78	P. 557. ² P. 37. ² Method 200.7. ⁴ 24

Table IB Notes

¹ "Methods for Analysis of Inorganic Substances in Water and Fluvial Sediments," U.S. Department of the Interior, U.S. Geological Survey, Open-File Report 78-679, or "Methods for Determination of Inorganic Substances in Water and Fluvial Sediments," NW Skougstad, et al. U.S. Geological Survey, Techniques of Water-Resources Investigation, Book 5, Chapter A1, 1979.

² "Official Methods of Analysis of the Association of Official Analytical Chemists" methods manual, 13th ed. (1980).

³ For the determination of total metals the sample is not filtered before processing. A digestion procedure is required to solubilize suspended material and to destroy possible organic-metal complexes. Two digestion procedures are given in "Methods for Chemical Analysis of Water and Wastes, 1979." One (§ 4.1.3), is a vigorous digestion using nitric acid. A less vigorous digestion using nitric and hydrochloric acid, (§ 4.1.4) is preferred, however, the analyst should be cautioned that this mild digestion may not suffice for all sample types. Particularly, if a colorimetric procedure is to be employed, it is necessary to ensure that all organo-metallic bonds be broken so that the metal is in a reactive state. In those situations, the vigorous digestion is to be preferred making certain that at no time does the sample go to dryness. Samples containing large amounts of organic materials would also benefit by this vigorous digestion. Use of the graphite furnace technique, inductively coupled plasma, as well as determinations for certain elements such as arsenic, the noble metals, mercury, selenium, and tellurium require a modified digestion and in all cases the method write-up should be consulted for specific instructions and/or cautions.

Note: If the digestion procedure for direct aspiration or graphite furnace atomic absorption analysis included in one of the other approved references is different than the above, the EPA procedure must be used.

Desolved metals are defined as those constituents which will pass through a 0.45 micron membrane filter. Following filtration of the sample, the referenced procedure for total metals must be followed. Sample digestion of the filtrate for dissolved metals, or digestion of the original sample solution for total metals may be omitted for AA (direct aspiration or graphite furnace) and ICP analyses provided the sample has a low COD and the filtrate meets the following criteria:

- is readily transparent;
- has no perceptible odor, and
- is free of particulate or suspended matter following acidification.

The full text of Method 200.7, "Inductively Coupled Plasma Atomic Emission Spectrometric Method for Trace Element Analysis of Water and Wastes," is given at Appendix C of this Part 136.

Manual distillation is not required if comparability data on representative effluent samples are on company file to show that the preliminary distillation step is not necessary; however, manual distillation will be required to resolve any controversies.

¹⁰ American Automated Electrode Method, Industrial Method Number 378-75WE, dated February 18, 1978, Technicon AutoAnalyzer II, Technicon Industrial Systems, Tarrytown, New York 10591.

¹¹ Carbonaceous biochemical oxygen demand (CBOD₅) must not be confused with the traditional BOD₅ test which measures "total BOD". The addition of the nitrification inhibitor is not a procedural option, but must be included to report the CBOD₅ parameter. A disclaimer whose permit requires reporting the traditional BOD₅ may not use a nitrification inhibitor in the procedure for reporting the results. Only when a disclaimer's permit specifically states CBOD₅ is required can the permittee report data obtained using the nitrification inhibitor.

¹² American National Standard on Photographic Processing Effluents, Apr. 2, 1975. Available from ANSI, 1430 Broadway, New York, NY 10018.

¹³ The use of normal and differential pulse voltage ramps to increase sensitivity and resolution is acceptable.

¹⁴ Chemical Oxygen Demand, Method 8000, Hach Handbook of Water Analysis, 1978, Hach Chemical Company, P.O. Box 388, Loveland, Colorado 80537.

¹⁵ COD Method, Oceanography International Corporation, 512 West Loop, P.O. Box 2980, College Station, Texas 77840.

¹⁶ The back titration method will be used to resolve controversy.

¹⁷ National Council of the Paper Industry for Air and Stream Improvement, Inc., Technical Bulletin 253, December 1971.

¹⁸ Copper, Bicinchoninate Method, Method 8508, Hach Handbook of Water Analysis, 1978, Hach Chemical Company, P.O. Box 388, Loveland, Colorado 80537.

¹⁹ After the manual distillation is completed, the auto-analyzer manifolds in EPA Methods 335.03 (Cyanide) or 420.2 (phenols) are simplified by connecting the re-sample line directly to the sampler. When using the manifold setup shown in Method 335, the buffer 6.2 should be replaced with the buffer 7.6 found in Method 335.2.

²⁰ Hydrogen Ion (pH) Automated Electrode Method, Industrial Method Number 378-75WA, October 1978, Technicon AutoAnalyzer II, Technicon Industrial Systems, Tarrytown, New York 10591.

²¹ Iron, 1,10-Phenanthroline Method, Method 8008, Hach Handbook of Water Analysis, 1978, Hach Chemical Company, P.O. Box 388, Loveland, Colorado 80537.

²² Manganese, Periodate Oxidation Method, Method 8034, Hach Handbook of Wastewater Analysis, 1978, pages 2-113 and 2-117, Hach Chemical Company, Loveland, Colorado 80537.

²³ Nitrogen, Nessler Method 8507, Hach Chemical Company, P.O. Box 388, Loveland, Colorado 80537.

²⁴ Goessens, D., Brown, E., "Methods for Analysis of Organic Substances in Water," U.S. Geological Survey, Techniques of Water-Resources Investigations, Book 5, Chapter A3, p.4 (1972).

²⁵ R.F. Anderson and R.G. Achman, "Direct Determination of Elemental Phosphorus by Gas-Liquid Chromatography," Journal of Chromatography, Vol. 47, No. 3, pp. 421-426, 1970.

²⁶ Recommended methods for the analysis of silver in industrial wastewaters at concentrations of 1 mg/L and above are inadequate where silver exists as an inorganic halide. Silver halides such as the bromide and chloride are relatively insoluble in reagents such as nitric acid but are readily soluble in an aqueous buffer of sodium thiosulfate and sodium hydroxide to a pH of 12. Therefore, for levels of silver above 1 mg/L, 20 mL of sample should be diluted to 100 mL by adding 40 mL each of 2 M Na₂S₂O₃ and 2M NaOH. Standards should be prepared in the same manner. For levels of silver below 1 mg/L the recommended method is satisfactory.

²⁷ Stevens, M.H., Ficks, J.F., and Smoot, G.F., "Water Temperature-Influential Factors, Field Measurement and Data Presentation," U.S. Geological Survey, Techniques of Water-Resources Investigations, Book 1, Chapter D1, 1975.

²⁸ Zinc, Zincon Method, Method 8008, Hach Handbook of Water Analysis, 1978, pages 2-231 and 2-333, Hach Chemical Company, Loveland, Colorado 80537.

²⁹ "Selected Analytical Methods Approved and Cited by the United States Environmental Protection Agency," Supplement to the Fifteenth Edition of Standard Methods for the Examination of Water and Wastewater (1981).

³⁰ The approved method is that cited in Standard Methods for the Examination of Water and Wastewater, 14th Edition. The colorimetric reaction is conducted at a pH of 10.0 ± 0.2. The approved methods are given on pp. 578-61 of the 14th Edition: Method 510A for distillation, Method 510B for the manual colorimetric procedure, or Method 510C for the manual spectrophotometric procedure.

³¹ ORION Research Instruction Manual, Residual Chlorine Electrode Model 97-70, 1977, Orion Research Incorporated, 840 Memorial Drive, Cambridge, Massachusetts 02138.

TABLE IC.—LIST OF APPROVED TEST PROCEDURES FOR NON-PESTICIDE ORGANIC COMPOUNDS

Parameter ¹	EPA Method Number ^{2,3}			Other
	GC	GC/MS	HPLC	
1. Acenaphthene	610	625, 1625	610	
2. Acenaphthylene	610	625, 1625	610	
3. Acridene	603	*624, 1624		
4. Acrylonitrile	603	*624, 1624		
5. Anthracene	610	625, 1625	610	
6. Benzene	602	624, 1624		
7. Benzidine		*625, 1625	605	Note 3, p. 1;
8. Benzo[a]anthracene	610	625, 1625	610	
9. Benzo[a]pyrene	610	625, 1625	610	
10. Benzo[b]fluoranthene	610	625, 1625	610	
11. Benzo[b]kynylene	610	625, 1625	610	
12. Benzo[b]fluoranthene	610	625, 1625	610	
13. Benzyl Chloride	610	625, 1625	610	Note 3, p. 130; Note 6, p. 5102.
14. Benzyl Butyl Phthalate	606	625, 1625		
15. Bis(2-chloroethoxy) methane	611	625, 1625		
16. Bis(2-chloroethyl) ether	611	625, 1625		
17. Bis(2-ethylhexyl) phthalate	606	625, 1625		
18. Bis(2-chloromethoxy) methane	601	624, 1624		
19. Bromobenzene	601	624, 1624		
20. Bromomethane	601	624, 1624		
21. 4-Bromophenyl phenyl ether	611	625, 1625		
22. Carbon tetrachloride	601	624, 1624		Note 3, p. 130;
23. 4-Chloro-3-methylphenol	604	625, 1625		
24. Chlorobenzene	601, 602	624, 1624		Note 3, p. 130;
25. Chloroethane	601	624, 1624		
26. 2-Chloroethyl vinyl ether	601	624, 1624		
27. Chloroform	601	624, 1624		Note 3, p. 130;
28. Chloromethane	601	624, 1624		
29. 2-Chloronaphthalene	612	625, 1625		
30. 2-Chlorophenol	604	625, 1625		
31. 4-Chlorophenyl phenyl ether	611	625, 1625		
32. Cryolite	610	625, 1625	610	
33. Dibenz[a,h]anthracene	610	625, 1625	610	
34. Dibenz[b,h]anthracene	601	624, 1624		
35. 1,3-Dichlorobenzene	601, 602, 612	624, 625, 1625		
36. 1,3-Dichlorobenzene	601, 602, 612	624, 625, 1625		
37. 1,4-Dichlorobenzene	601, 602, 612	625, 1624, 1625		
38. 2,3-Dichlorobenzidine		625, 1625	606	
39. Dichlorodifluoromethane	601			
40. 1,1-Dichloroethane	601	624, 1624		
41. 1,2-Dichloroethane	601	624, 1624		
42. 1,1-Dichloroethene	601	624, 1624		
43. trans-1,2-Dichloroethene	601	624, 1624		
44. 2,4-Dichlorophenol	604	625, 1625		
45. 1,3-Dichloropropane	601	624, 1624		
46. cis-1,3-Dichloropropane	601	624, 1624		
47. trans-1,3-Dichloropropane	601	624, 1624		
48. Diethyl phthalate	606	625, 1625		

TABLE IC.—LIST OF APPROVED TEST PROCEDURES FOR NON-PESTICIDE ORGANIC COMPOUNDS—Continued

Parameter ¹	EPA Method Number ^{2,3}			Other
	GC	GC/MS	HPLC	
48. 2,4-Dimethylphenol	804	625, 1625		Note 3, p. 130; Note 6, p. 5102.
49. Dimethyl phthalate	806	625, 1625		
50. Di-n-butyl phthalate	806	625, 1625		
51. Di-n-octyl phthalate	806	625, 1625		
52. 2,4-Dichlorophenol	804	625, 1625		
53. 2,4-Dichlorobenzene	808	625, 1625		
54. 2,6-Dichlorobenzene	808	625, 1625		
55. Epichlorohydrin				
57. Ethylbenzene	802	624, 1624		Note 3, p. 130;
58. Fluoranthene	810	625, 1625	610	
59. Fluorene	810	625, 1625	610	
60. Hexachlorobenzene	812	625, 1625		
61. Hexachlorobutadiene	812	625, 1625		
62. Hexachlorocyclopentadiene	812	625, 1625		
63. Hexachloroethane	812	625, 1625		
64. Menthyl 2,3-cylopyrene	810	625, 1625	610	
65. Isophorone	808	625, 1625		Note 3, p. 130;
66. Methylene Chloride	801	624, 1624		
67. 2-Methyl-4,6-Dinitrophenol	804	625, 1625		
68. Naphthalene	810	625, 1625		
69. Nitrobenzene	808	625, 1625		
70. 2-Nitrophenol	804	625, 1625		
71. 4-Nitrophenol	804	625, 1625		
72. N-Nitrosodimethylamine	807	625, 1625		
73. N-Nitroso-n-propylamine	807	625, 1625		Note 3, p. 43; Note 3, p. 43; Note 3, p. 43; Note 3, p. 43; Note 3, p. 43; Note 3, p. 43; Note 3, p. 43; Note 3, p. 140;
74. N-Nitrosodiphenylamine	807	625, 1625		
75. 2,3-dinitro-1-chloropropane	811	625, 1625		
76. PCB-1016	808		625	
77. PCB-1221	808		625	
78. PCB-1232	808		625	
79. PCB-1242	808		625	
80. PCB-1248	808		625	
81. PCB-1254	808		625	
82. PCB-1280	808		625	
83. Pentachlorophenol	804	625, 1625		Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130;
84. Phenanthrene	810	625, 1625	610	
85. Phenol	804	625, 1625		
86. Pyrene	810	625, 1625	610	
87. 2,3,7,8-Tetrachlorodibenzo-p-dioxin		813		
88. 1,1,2,3-Tetrachloronaphthalene	801	624, 1624		
89. Tetrachloroethene	801	624, 1624		
90. Toluene	802	624, 1624		
91. 1,2,4-Trichlorobenzene	812	625, 1625		Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130; Note 3, p. 130;
92. 1,1,1-Trichloroethane	801	624, 1624		
93. 1,1,2-Trichloroethane	801	624, 1624		
94. Trichloroethene	801	624, 1624		
95. Trichlorofluoromethane	801	624		
96. 2,4,6-Trichlorophenol	804	625, 1625		
97. Vinyl Chloride	801	624, 1624		

Table IC Notes

¹ All parameters are expressed in micrograms per liter (µg/L).² The full text of Methods 801-813, 624, 625, 1624, and 1625, are given at Appendix A, "Test Procedures for Analysis of Organic Pollutants," of this Part 136. The standardized test procedure to be used to determine the method detection limit (MDL) for these test procedures is given at Appendix B, "Definition and Procedure for the Determination of the Method Detection Limit," of this Part 136.³ Methods for Benzidine: Chlorinated Organic Compounds, Pentachlorophenol and Pesticides in Water and Wastewater," U.S. Environmental Protection Agency, September, 1978. Method 624 may be extended to screen samples for Acroten and Acrylonitrile. However, when they are known to be present, the preferred method for these two compounds is Method 803 or Method 1624.⁴ Method 625 may be extended to include benzidine, hexachlorocyclopentadiene, N-nitrosodimethylamine, and N-nitrosodiphenylamine. However, when they are known to be present, Methods 805, 807, and 812, or Method 1625, are preferred methods for these compounds.⁵ 625, Screening only.⁶ Selected Analytical Methods Approved and Cited by the United States Environmental Protection Agency," Supplement to the Fifteenth Edition of Standard Methods for the Examination of Water and Wastewater (1981).⁷ Each analyst must make an initial, one-time, demonstration of their ability to generate acceptable precision and accuracy with Methods 801-813, 624, 625, 1624, and 1625 (See Appendix A of this Part 136) in accordance with procedure each in section 8.2 of each of these Methods. Additionally, each laboratory, on an on-going basis must spike and analyze 10% (5% for Methods 804 and 825 and 100% for methods 1624 and 1625) of all samples to monitor and evaluate laboratory data quality in accordance with sections 8.3 and 8.4 of these Methods. When the recovery of any parameter falls outside the warning limits, the analytical results for that parameter in the unspiked sample are suspect and cannot be reported to demonstrate regulatory compliance.

Note.—These warning limits are promulgated as an "interim final action with a request for comments."

TABLE ID.—LIST OF APPROVED TEST PROCEDURES FOR PESTICIDES¹

Parameter µg/L	Method	EPA ^{2,3}	Standard Methods 15th Ed	ASTM	Other
1. Aldrin	GC	808	508A	D3086	Note 3, p. 7; Note 4, p. 30.
	GC/MS	625			
2. Atrazine	GC				Note 3, p. 83; Note 6, p. 508.
3. Avenacarb	TLC				
4. Atrazine	GC				Note 3, p. 84; Note 6, p. 518.
5. Atrazine	GC				
6. Atrazine methyl	GC				Note 3, p. 83; Note 6, p. 508.
7. Barban	TLC				
8. o-BHC	GC	808	508A	D3086	Note 3, p. 25; Note 6, p. 551.
	GC/MS	625			
9. p-BHC	GC	808		D3086	Note 3, p. 104; Note 6, p. 564.
	GC/MS	625			
10. t-BHC	GC	808		D3086	Note 3, p. 7

TABLE ID.—LIST OF APPROVED TEST PROCEDURES FOR PESTICIDES¹—Continued

Parameter µg/L	Method	EPA ²	Standard Methods 15th Ed	ASTM	Other
11. γ-BHC (Lindane)	GC/MS GC GC/MS GC	625 608 625	508A	D3086	Note 3, p. 7, Note 4, p. 30.
12. Captafen	TLC		508A		Note 3, p. 7
13. Carbofuryl	GC				Note 3, p. 94, Note 6, p. S80.
14. Carbofenthothion	GC				Note 4, p. 30, Note 6, p. S73.
15. Chlorobenzene	GC GC-MS	608 625	508A	D3086	Note 3, p. 7.
16. Chlorobutanol	TLC				Note 3, p. 104, Note 6, p. S84.
17. 2,4-D	GC		508B		Note 3, p. 115, Note 4, p. 35.
18. 4,4'-DDD	GC GC-MS	608 625	508A	D3086	Note 3, p. 7, Note 4, p. 30.
19. 4,4'-DDE	GC GC/MS	608 625	508A	D3086	Note 3, p. 7, Note 4, p. 30.
20. 4,4'-DDT	GC GC/MS	608 625	508A	D3086	Note 3, p. 7, Note 4, p. 30.
21. Dactinon-O	GC				Note 3, p. 25, Note 6, p. S51.
22. Dactinon-S	GC				Note 3, p. 25, Note 6, p. S51
23. Dactinon	GC				Note 3, p. 25, Note 4, p. 30, Note 6, p. S51
24. Dactinon	(M)				Note 1, p. 115
25. Dichlorophenol	GC				Note 4, p. 30, Note 6, p. S73
26. Dichlorophenol	GC		508A		Note 3, p. 7
27. Dieldrin	GC			D3086	Note 3, p. 7
28. Dieldrin	GC GC/MS	608 625	508A		Note 3, p. 7, Note 4, p. 30.
29. Dieldrin	GC				Note 4, p. 30, Note 6, p. S73.
30. Dieldrin	GC				Note 3, p. 7, Note 6, p. S51.
31. Dieldrin	TLC				Note 3, p. 104, Note 6, p. S64.
32. Endosulfan I	GC GC/MS	608 625	508A	D3086	Note 3, p. 7.
33. Endosulfan II	GC GC/MS	608 625	508A	D3086	Note 3, p. 7.
34. Endosulfan sulfate	GC GC/MS	608 625			
35. Endrin	GC GC/MS	608 625	508A	D3086	Note 3, p. 7, Note 4, p. 30.
36. Endrin aldehyde	GC GC/MS	608 625			
37. Ethion	GC				Note 4, p. 30, Note 6, p. S73.
38. Fenitrothion	TLC				Note 3, p. 104, Note 6, p. S64.
39. Fenitrothion-TCA	TLC				Note 3, p. 104, Note 6, p. S64.
40. Heptachlor	GC GC/MS	608 625	508A	D3086	Note 3, p. 7, Note 4, p. 30.
41. Heptachlor epoxide	GC	608	508A	D3086	Note 3, p. 7, Note 4, p. 30, Note 6, p. S73.
42. Isodrin	GC/MS	625			
43. Lincosin	GC				Note 4, p. 30, Note 6, p. S73.
44. Malathion	TLC GC		508A		Note 3, p. 104, Note 6, p. S64.
45. Malathion	GC				Note 3, p. 25, Note 4, p. 30, Note 6, p. S51.
46. Malathion	TLC				Note 3, p. 94, Note 6, p. S80.
47. Malathion	GC		508A	D3086	Note 3, p. 7, Note 4, p. 30.
48. Malathion	TLC				Note 3, p. 94, Note 6, p. S80.
49. Malathion	GC		508A		Note 3, p. 7.
50. Malathion	TLC				Note 3, p. 104, Note 6, p. S84.
51. Malathion-TCA	TLC				Note 3, p. 104, Note 6, p. S84.
52. Malathion	TLC				Note 3, p. 104, Note 6, p. S64.
53. Parathion methyl	GC		508A		Note 3, p. 25, Note 4, p. 30.
54. Parathion ethyl	GC		508A		Note 3, p. 25.
55. PCNB	GC		508A		Note 3, p. 7.
56. Permethrin	GC			D3086	
57. Permethrin	GC				Note 3, p. 83, Note 6, p. S68.
58. Permethrin	GC				Note 3, p. 83, Note 6, p. S68.
59. Permethrin	GC				Note 3, p. 83, Note 6, p. S68.
60. Permethrin	TLC				Note 3, p. 104, Note 6, p. S64.
61. Permethrin	TLC				Note 3, p. 84, Note 6, p. S80.
62. Sebacic acid	TLC				Note 3, p. 83, Note 6, p. S68.
63. Sebacic acid	TLC				Note 3, p. 104, Note 6, p. S64.
64. Sebacic acid	GC				Note 3, p. 83, Note 6, p. S68.
65. Sebacic acid	GC		508A		Note 3, p. 7.
66. Sebacic acid	TLC				Note 3, p. 104, Note 6, p. S64.
67. 2,4,5-T	GC		508B		Note 3, p. 115, Note 4, p. 35.
68. 2,4,5-TP (Silver)	GC		508B		Note 3, p. 115
69. Terbufosfate	GC				Note 3, p. 83, Note 6, p. S68
70. Terbufosfate	GC GC/MS GC	608 625	508A	D3086	Note 3, p. 7, Note 4, p. 30

Table ID Notes

- ¹ Pesticides are listed in this table by common name for the convenience of the reader. Additional pesticides may be found under Table IC, where entries are listed by chemical name.
- ² The full text of methods 608 and 625 are given at Appendix A, "Test Procedures for Analysis of Organic Pollutants," of the Part 136. The standardized test procedure to be used to determine the method detection limit (MDL) for these test procedures is given at Appendix B, "Detection and Procedure for the Determination of the Method Detection Limit," of the Part 136.
- ³ Methods for Benzidine, Chlorinated Organic Compounds, Pentachlorophenol and Pesticides in Water and Wastewater, U.S. Environmental Protection Agency, September, 1978. This EPA publication includes thin-layer chromatography (TLC) methods.
- ⁴ Methods for Analysis of Organic Substances in Water, U.S. Geological Survey, Techniques of Water-Resources Investigations, Book 5, Chapter A3 (1972).
- ⁵ This method may be extended to include α-BHC, β-BHC, endosulfan I, endosulfan II, and endrin. However, when they are known to exist, Method 608 is the preferred method.
- ⁶ Standard Analytical Methods Approved and Used by the United States Environmental Protection Agency, "Supplement to the Fifteenth Edition of Standard Methods for the Examination of Water and Wastewater (1981).
- ⁷ Each analyst must make an initial, one-time, demonstration of their ability to generate acceptable precision and accuracy with Methods 608 and 625 (See Appendix A of the Part 136) in accordance with procedures given in Section 8.2 of each of these methods. Additionally, each laboratory, on an on-going basis, must spike and analyze 10% of all samples analyzed with

Method 808 or 5% of all samples analyzed with Method 825 to monitor and evaluate laboratory data quality in accordance with Sections 8.3 and 8.4 of these methods. When the recovery of any parameter falls outside the warning limits, the analytical results for that parameter in the unspiked sample are suspect and cannot be reported to demonstrate regulatory compliance.

Note.—These warning limits are promulgated as an "interim final action with a request for comments."

TABLE IE.—LIST OF APPROVED RADIOLOGICAL TEST PROCEDURES

Parameter and units	Methods	FPA*	Reference (method No. or page)		
			Standard Methods 15th Ed	ASTM	USGS*
1. Alpha-Test, μCi per liter	Proportional or scintillation counter	900 0	703	D1943-66	pp 75 and 78. ^b
2. Alpha-Counting error, μCi per liter	Proportional or scintillation counter	Appendix B	703	D1943-66	p 79.
3. Alpha-Counting error, μCi per liter	Proportional counter	900 0	703	D1890-66	pp 75 and 78. ^b
4. Beta-Counting error, μCi per liter	Proportional counter	Appendix B	703	D1890-66	p 79.
5. (a) Radium-Total, μCi per liter	Proportional counter	903 0	705	D2460-70	
(b) mCi , μCi per liter	Scintillation counter	903 1	705	D2454-79	p 81.

Table IE Notes

* "Prescribed Procedures for Measurement of Radioactivity in Drinking Water." EPA-600/4-80-032 (1980 update), U.S. Environmental Protection Agency, August 1980.

^b Fawcett, M.J. and Brown, Eugene, "Selected Methods of the U.S. Geological Survey of Analysis of Wastewaters." U.S. Geological Survey, Open-File Report 78-177 (1978).

^c The method found on p. 75 measures only the dissolved portion while the method on p. 78 measures only the suspended portion. Therefore, the two results must be added to obtain the "total."

5. In § 136.3, paragraph (a) is revised to show that the full text of approved test procedures have been incorporated by reference, into the regulation to read as follows:

§ 136.3 Identification of test procedures.

(a) Parameters or pollutants, for which methods are approved, are listed together with test procedure descriptions and references in Tables IA, IB, IC, ID, and IE. The full text of the referenced test procedures are incorporated by reference into Tables IA, IB, IC, ID, and IE. The references and the sources from which they are available are given in paragraph (b) of this section. These test procedures are incorporated as they exist on the day of approval and a notice of any change in these test procedures will be published

in the Federal Register. The discharge parameter values for which reports are required must be determined by one of the standard analytical test procedures incorporated by reference and described in Tables IA, IB, IC, ID, and IE, or by any alternate test procedure which has been approved by the Administrator under the provisions of paragraph (d) of this section and sections 136.4 and 136.5 of this Part 136. Under certain circumstances (§§ 136.3 (b) or (c) or 40 CFR Part 401.13) other test procedures may be used that may be more advantageous when such other test procedures have been previously approved by the Regional Administrator of the Region in which the discharge will occur, and providing the Director of the State in which such discharge will occur

does not object to the use of such alternate test procedure.

6. In § 136.3, paragraphs (b) and (c) are redesignated as (c) and (d) and a new paragraph (b) is added to itemize the references which are "incorporated by reference" and to identify the sources from which they may be obtained. As added, the new paragraph (b) reads as follows:

§ 136.3 Identification of test procedures.

(b) The full texts of the methods from the following references which are cited in Tables IA, IB, IC, ID, and IE are incorporated by reference into this regulation and may be obtained from the sources identified. All costs cited are subject to change and must be verified from the indicated sources.

REFERENCES, SOURCES, AND COSTS

Table	Parameters	Reference, source and cost
IA—EPA	1-5	"Microbiological Methods for Monitoring the Environment, Water and Wastes." United States Environmental Protection Agency, EPA-600/8-78-017, 1978. Available from ORD Publications, CERL, U.S. Environmental Protection Agency, Cincinnati, Ohio 45268.
IA—Standard Methods	1-5	Standard Methods for the Examination of Water and Wastewater, Joint Editorial Board, American Public Health Association, American Water Works Association, and Water Pollution Control Federation 15th Edition, 1981. Available from: American Public Health Association, 1015 Fifteenth Street, N.W., Washington, D.C. 20036. Cost \$50.00 including the Supplement to the Fifteenth Edition 1981, 14th Edition.
IB—Standard Methods	1-10, 12-46, 50-75	
IC—Standard Methods	1, 8, 11, 12, 15, 17-20, 26, 28, 32, 33, 35, 40, 41, 44, 46, 48, 52-54, 64, 66, 67, 69, 70	
IE—Standard Methods	1-5	
IB—Standard Methods	48	
IC—Other (Standard Methods Supplement)	11, 47	
ID—Other (Standard Methods Supplement)	13, 56	
IE—Other (Standard Methods Supplement)	2-7, 13, 14, 16, 21-23, 25, 29-31, 37, 38, 39, 41, 45, 47, 49, 50, 51, 56-63, 65, 68	"Selected Analytical Methods approved and Used by the United States Environmental Protection Agency." Supplement to the 15th Edition of Standard Methods for the Examination of Water and Wastewater (1981). Available from: American Public Health Association, 1015 Fifteenth Street, N.W., Washington, D.C. 20036. Cost included with the 15th Edition of Standard Methods for the Examination of Water and Wastewater.
IA—U.S. Geological Survey (USGS)	1, 3, 5	"Methods for Collection and Analysis of Aquatic Biological and Microbiological Samples," edited by P.E. Greeson, T.A. Ehle, G.A. Ivem, B.W. Lum, and K.V. Slack. U.S. Geological Survey, Techniques of Water-Resources Investigation (USGS TWRI), Book 5, Chapter A4 (1977). Revised edition, 332 pages. Available from U.S. Geological Survey, Branch of Distribution, 1200 South Eads Street, Arlington, VA 22202. (Authorized agent of the Superintendent of Documents, Government Printing Office.) Cost: \$9.25. Prices are subject to change.
IB—EPA	1-13, 15-48, 50-75	"Methods for Chemical Analysis of Water and Wastes", EPA-600/4-79-020 United States Environmental Protection Agency, March, 1979. Available from ORD Publications, CERL, U.S. Environmental Protection Agency Cincinnati, Ohio 45268.
IB—ASTM	1, 2, 4, 6, 8, 11-13, 15-17, 19, 20, 22-25, 27, 29, 30-35, 37-40, 42-44, 46, 48, 50, 52, 60, 61, 63-65, 67, 68, 73-75	"Annual Book of Standards, Part 31, Water", American Society for Testing and Materials, 1980. Available from American Society for Testing and Materials, 1916 Race Street, Philadelphia, PA 19103. Cost available from publisher.
ID—ASTM	1, 8-11, 13, 18-20, 27, 32, 33, 35, 40, 41, 46, 55, 69	
IE—ASTM	1-5	

REFERENCES, SOURCES, AND COSTS—Continued

Table	Parameters	Reference, source and cost
USGS	2, 3, 4, 6-13, 15, 16, 18-23, 25, 27, 28, 30-40, 43, 44, 46, 50, 52-55, 57, 60-66, 71, 73, 75.	"Methods for determination of inorganic substances in water and fluvial sediments," N.W. Skougstad and others, editors USGS-TWRI Book 5, Chapter A1, 1979. \$10.00. Available from U.S. Geological Survey, Branch of Distribution, 1200 South Eads Street, Arlington, VA 22202 (Authorized agent of the Superintendent of Documents, Government Printing Office). Prices are subject to change.
Other (AOAC)	2, 4, 6, 12, 15, 16, 19, 22, 30-35, 38, 42-44, 46, 50, 52, 62-65, 75.	Official Methods of Analysis of the Association of Official Analytical Chemists, methods manual, 13th Edition (1980). Price: \$78.00. Available from: The Association of Official Analytical Chemists, 1111 N. 19th St., Suite 210, Arlington, VA 22209.
Other (ANSI)	8, 12, 15, 20, 22, 23, 38, 62, 75.	"American National Standard on Photographic Processing Effluents," April 2, 1975. Available from American National Standards Institute, 1430 Broadway, New York, New York 10018.
Other (EPA)	3, 5-8, 10, 12, 13, 19, 20, 22, 27, 30, 32-34, 36, 37, 42, 43, 47, 74, 75.	The full text of the inductively coupled plasma optical emission spectroscopic test procedure, Method 200.7, is printed in Appendix C of this Part 136.
Other	71	"An Investigation of Improved Procedures for Measurement of Mill Effluent and Heavy Metals in Mill Aqueous Effluents," M. A. J. (unpublished), 1971. Available from National Council of the Paper Industry for Air and Waste Improvements, Inc., 2701 Madison Avenue, Cincinnati, Ohio 45226.
Other	4	Ammonia, Automated Electrode Method, Industrial Method Number 379-75WE, dated February 19, 1976, Technicon AutoAnalyzer II Method and price available from Technicon Industrial Systems, Tarrytown, New York 10591.
Other	15	Chemical Oxygen Demand, Method 8000, Hach Handbook of Water Analysis, 1979. Method and price available from Hach Chemical Company, P.O. Box 389, Loveland, Colorado 80537.
Other	15	DOC Chemical Oxygen Demand Method, Method and price available from Oceanography International Corporation, 512 West Loop, P.O. Box 2960, College Station, Texas 77840.
Other	17	ORION Research Instruction Manual, Residual Chlorine Electrode Model 97-78, 1977. Method and price available from Orion Research Incorporated, 840 Memorial Drive, Cambridge, Massachusetts 02138.
Other	22	Bicinchonate Method for Copper, Method 8506, Hach Handbook of Water Analysis, 1979. Method and price available from Hach Chemical Company, P.O. Box 389, Loveland, Colorado 80537.
Other	28	Hydrogen Ion (pH) Automated Electrode Method, Industrial Method Number 378-75WA, October 1976, Technicon AutoAnalyzer II Method and Price available from Technicon Industrial Systems, Tarrytown, New York 10591.
Other	30	1, 10-Phenanthroline Method for Iron, Hach Method 8008. Method and price available from Hach Chemical Company, P.O. Box 389, Loveland, Colorado 80537.
Other	34	Periodate Oxidation Method for Manganese, Method 8034, Hach Handbook for Water Analysis, 1979. Method and Price available from Hach Chemical Company, P.O. Box 389, Loveland, Colorado 80537.
Other	40	Nitrite Nitrogen, Hach Method 8507. Method and price available from Hach Chemical Company, P.O. Box 389, Loveland, Colorado 80537.
Other	75	Zincin Method for Zinc, Method 8009, Hach Handbook for Water Analysis, 1979. Method and price available from Hach Chemical Company, P.O. Box 389, Loveland, Colorado 80537.
Other	49	"Direct Determination of Elemental Phosphorus by Gas-Liquid Chromatography," by R.F. Addison and R.G. Ackman, Journal of Chromatography, Volume 47, No. 3, pp. 421-426, 1970. Available in most public libraries. Back volumes of the Journal of Chromatography are available from Elsevier/North-Holland, Inc., Journal Information Centre, 52 Vanderbilt Avenue, New York, NY 10164. Cost available from publisher.
Other (USGS)	69	"Water temperature-influential factors, field measurement, and data presentation," by H.H. Stevens, Jr., J. Ficke, and G.F. Smoot: USGS-TWRI Book 1, Chapter D1, 1975. 65 pages, \$1.60. Available from U.S. Geological Survey, Branch of Distribution, 1200 South Eads Street, Arlington, VA 22202. Prices are subject to change.
Other (USGS) ID-Other (USGS)	42 1, 11, 14, 17-20, 23, 25, 28, 29, 35, 37, 40-42, 44, 46, 52, 66, 68	"Methods for analysis of organic substances in water," by D. F. Goertitz and Eugene Brown, USGS-TWRI Book 5, Chapter A3, 1972, 40 pages, \$3.00. Available from U.S. Geological Survey, Branch of Distribution, 1200 South Eads Street, Arlington, VA 22202. Prices are subject to change.
IC-EPA ID-EPA	1-12, 14-55, 57-67 1, 8-11, 15, 18, 19, 20, 28, 32-36, 40, 41, 68	The full texts of Methods 601-613, 624, 625, 1624, and 1625 are printed in appendix A of this Part 136. The full text for determining the method detection limit when using the test procedures is given in Appendix B of this Part 136.
IC-Other (EPA) ID-Other	7, 13, 22, 24, 27, 56, 66, 76-83, 86, 88, 91, 93 1-8, 11-13, 15-24, 26, 28, 30-33, 35, 38-41, 43-54, 56-70	"Methods for Benzidine, Chlorinated Organic Compounds, Pentachlorophenol and Pesticides in Water and Wastewater," Environmental Monitoring and Support Laboratory, United States Environmental Protection Agency, Cincinnati, Ohio 1978. Available from ORD Publications, CERL, U.S. Environmental Protection Agency, Cincinnati, Ohio 45268.
IE-EPA	1-5	"Prescribed Procedures for Measurement of Radioactivity in Drinking Water," EPA-800/4-80-032 (1980 Update), United States Environmental Protection Agency, 1980. Available from ORD Publications, CERL, U.S. Environmental Protection Agency, Cincinnati, Ohio 45268.
IE-USGS	1-5	"Selected Methods of the U.S. Geological Survey of Analysis of Wastewaters," by M.J. Fishman and Eugene Brown, U.S. Geological Survey Open File Report 76-77 (1976) \$13.50. Available from: U.S. Geological Survey, Branch Distribution, 1200 South Eads Street, Arlington, VA 22202.

The full texts of all the test procedures cited are available for inspection at the Office of the Federal Register Information Center, Room 8301, 1110 L Street, N.W., Washington, D.C. 20408.

7. In section 136.3 a new paragraph (e) is added together with a new Table II entitled, "Table II, Required Containers, Preservation Techniques, and Holding Times," to read as follows:

§ 136.3 Identification of test procedures.

(e) Sample preservation procedures, container materials, and maximum allowable holding times for parameters cited in Tables IA, IB, IC, ID, and IE are

prescribed in Table II. Any person may apply for a variance from the prescribed preservation techniques, container materials, and maximum holding times applicable to samples taken from a specific discharge. Applications for variances may be made by letters to the Regional Administrator in the Region in which the discharge will occur. Sufficient data should be provided to

assure such variance does not adversely affect the integrity of the sample. Such data will be forwarded by the Regional Administrator to the Director of the Environmental Monitoring and Support Laboratory in Cincinnati, Ohio for technical review and recommendations for action on the variance application. Upon receipt of the recommendations from the Director of the Environmental

Monitoring and Support Laboratory, the Regional Administrator may grant a variance applicable to the specific discharge to the applicant. A decision to approve or deny a variance will be made within 90 days of receipt of the application by the Regional Administrator.

TABLE II.—REQUIRED CONTAINERS, PRESERVATION TECHNIQUES, AND HOLDING TIMES

Parameter No./name	Container ¹	Preservation ^{1,2}	Maximum holding time ³
Table IA—Bacterial Tests:			
1-4. Coliform, fecal and total	P, G	Cool, 4°C, 0.008% Na ₂ S ₂ O ₅ ⁴	6 hours.
5. Fecal streptococci	P, G	do	Do.
Table IB—Inorganic Tests:			
1. Acidity	P, G	Cool, 4°C	14 days.
2. Alkalinity	P, G	do	Do.
4. Ammonia	P, G	Cool, 4°C, H ₂ SO ₄ to pH < 2	28 days.
9. Biochemical oxygen demand	P, G	Cool, 4°C	48 hours.
11. Bromide	P, G	None required	28 days.
14. Biochemical oxygen demand, carbonaceous	P, G	Cool, 4°C	48 hours.
15. Chemical oxygen demand	P, G	Cool, 4°C, H ₂ SO ₄ to pH < 2	28 days.
16. Chloride	P, G	None required	Do.
17. Chlorine, total residual	P, G	do	Analyze immediately.
21. Color	P, G	Cool, 4°C	48 hours.
23-24. Cyanide, total and amenable to chlorination	P, G	Cool, 4°C, NaOH to pH > 12, 0.5g ascorbic acid ⁵	14 days ⁶
25. Fluoride	P	None required	28 days.
27. Hardness	P, G	HNO ₃ to pH < 2, H ₂ SO ₄ to pH < 2	6 months.
28. Hydrogen ion (pH)	P, G	None required	Analyze immediately.
31, 43. Kjeldahl and organic nitrogen	P, G	Cool, 4°C, H ₂ SO ₄ to pH < 2	28 days.
Metals⁷:			
18. Chromium VI	P, G	Cool, 4°C	24 hours.
25. Mercury	P, G	HNO ₃ to pH < 2	28 days.
3, 5-8, 10, 12, 13, 19, 20, 22, 26, 29, 30, 32-34, 36, 37, 45, 47, 51, 52, 58-60, 62, 63, 70-72, 74, 75. Metals, except chromium VI and mercury	P, G	do	6 months.
38. Nitrate	P, G	Cool, 4°C	48 hours.
39. Nitrate-nitrite	P, G	Cool, 4°C, H ₂ SO ₄ to pH < 2	28 days.
48. Nitrite	P, G	Cool, 4°C	48 hours.
41. Oil and grease	G	Cool, 4°C, H ₂ SO ₄ to pH < 2	28 days.
42. Organic carbon	P, G	Cool, 4°C, HCl or H ₂ SO ₄ to pH < 2	Do.
44. Orthophosphate	P, G	Fair immediately. Cool, 4°C	48 hours.
46. Oxygen, Dissolved Probe	G. Bottle and top	None required	Analyze immediately.
47. Winstar	do	Fix on site and store in dark	8 hours.
48. Phenols	G only	Cool, 4°C, H ₂ SO ₄ to pH < 2	28 days.
49. Phosphorus (elemental)	G	Cool, 4°C	48 hours.
50. Phosphorus, total	P, G	Cool, 4°C, H ₂ SO ₄ to pH < 2	28 days.
53. Residue, total	P, G	Cool, 4°C	7 days.
54. Residue, Filterable	P, G	do	48 hours.
55. Residue, Nonfilterable (TSS)	P, G	do	7 days.
56. Residue, Settleable	P, G	do	48 hours.
57. Residue, volatile	P, G	do	7 days.
61. Silica	P	do	28 days.
64. Specific conductance	P, G	do	Do.
65. Sulfate	P, G	do	Do.
66. Sulfide	P, G	Cool, 4°C add zinc acetate plus sodium hydroxide to pH > 9	7 days.
67. Sulfite	P, G	None required	Analyze immediately.
68. Surfactants	P, G	Cool, 4°C	48 hours.
69. Temperature	P, G	None required	Analyze.
73. Turbidity	P, G	Cool, 4°C	48 hours.
Table IC—Organic Tests⁸:			
13, 18-20, 22, 24-26, 34-37, 39-43, 45-47, 56, 66, 68, 69, 92-95, 97. Purgeable Halocarbons	G, Teflon-lined septum	Cool, 4°C, 0.008% Na ₂ S ₂ O ₅ ⁴	14 days.
6, 57, 90. Purgeable aromatic hydrocarbons	do	Cool, 4°C, 0.008% Na ₂ S ₂ O ₅ ⁴ , HCl to pH 0 ⁹	No
3, 4. Acrolein and acrylonitrile	do	Cool, 4°C, 0.008% Na ₂ S ₂ O ₅ ⁴ , Adjust pH to 4-5 ¹⁰	Do
23, 30, 44, 49, 53, 67, 70, 71, 83, 85, 96. Phenols ¹¹	G, Teflon-lined cap	Cool, 4°C, 0.008% Na ₂ S ₂ O ₅ ⁴	7 days until extraction, 40 days after extraction.
7, 38. Benzidines ¹²	do	do	7 days until extraction ¹³
14, 17, 48, 50-52. Phenolate esters ¹⁴	do	Cool, 4°C	7 days until extraction, 40 days after extraction.
72-74. Neuroamines ¹⁵	do	Cool, 4°C store in dark, 0.008% Na ₂ S ₂ O ₅ ⁴	Do
76-82. PCBs ¹⁶ acrylonitrile	do	Cool, 4°C	Do
54, 55, 65, 69. Neurotoxins and isophorone ¹⁷	do	Cool, 4°C, 0.008% Na ₂ S ₂ O ₅ ⁴ store in dark	Do
1, 2, 5, 8-12, 32, 33, 58, 59, 64, 68, 84, 86. Polynuclear aromatic hydrocarbons ¹⁸	do	do	Do
15, 16, 21, 31, 75. Halothanes ¹⁹	do	Cool, 4°C, 0.008% Na ₂ S ₂ O ₅ ⁴	Do
29, 35-37, 60-63, 91. Chlorinated hydrocarbons ²⁰	do	Cool, 4°C	Do
87. TCDD ²¹	do	Cool, 4°C, 0.008% Na ₂ S ₂ O ₅ ⁴	Do
Table ID—Pesticides Tests:			
1-70. Pesticides ²²	do	Cool, 4°C, pH 5-9 ²³	Do
Table IE—Radiological Tests:			
1-5. Alpha, beta and radium	P, G	HNO ₃ to pH < 2	6 months.

Table II Notes

¹ Polyethylene (P) or Glass (G)

* Sample preservation should be performed immediately upon sample collection. For composite chemical samples each aliquot should be preserved at the time of collection. When use of an automated sampler makes it impossible to preserve each aliquot, then chemical samples may be preserved by maintaining at 4°C until compositing and sample splitting is completed.

* When any sample is to be shipped by common carrier or sent through the United States Mails, it must comply with the Department of Transportation Hazardous Materials Regulations (49 CFR Part 172). The person offering such material for transportation is responsible for ensuring such compliance. For the preservation requirements of Table II, the Office of Hazardous Materials, Materials Transportation Bureau, Department of Transportation has determined that the Hazardous Materials Regulations do not apply to the following materials: Hydrochloric acid (HCl) in water solutions at concentrations of 0.04% by weight or less (pH about 1.96 or greater); Nitric acid (HNO₃) in water solutions at concentrations of 0.15% by weight or less (pH about 1.82 or greater); Sulfuric acid (H₂SO₄) in water solutions at concentrations of 0.35% by weight or less (pH about 1.15 or greater); and Sodium hydroxide (NaOH) in water solutions at concentrations of 0.080% by weight or less (pH about 12.30 or less).

* Samples should be analyzed as soon as possible after collection. The times listed are the maximum times that samples may be held before analysis and still be considered valid. Samples may be held for longer periods only if the permittee, or monitoring laboratory, has data on file to show that the specific types of samples under study are stable for the longer time, and has received a variance from the Regional Administrator under § 136.3(e). Some samples may not be stable for the maximum time period given in the table. A permittee, or monitoring laboratory, is obligated to hold the sample for a shorter time if knowledge exists to show that this is necessary to maintain sample stability. See § 136.3(e) for details.

* Should only be used in the presence of residual chlorine.

* Maximum holding time is 24 hours when sulfide is present. Optionally all samples may be tested with lead acetate paper before pH adjustments in order to determine if sulfide is present. If sulfide is present, it can be removed by the addition of calcium nitrate powder until a negative spot test is obtained. The sample is filtered and then NaOH is added to pH 12.

* Samples should be filtered immediately on-site before adding preservative for dissolved metals.

* Guidance applies to samples to be analyzed by GC, LC, or GC/MS for specific compounds.

* Samples receiving no pH adjustment must be analyzed within seven days of sampling.

* The pH adjustment is not required if acetone will not be measured. Samples for acetone receiving no pH adjustment must be analyzed within 3 days of sampling.

* When the extractable analyses of concern fall within a single chemical category, the specified preservative and maximum holding times should be observed for optimum safeguard of sample integrity. When the analyses of concern fall within two or more chemical categories, the sample may be preserved by cooling to 4°C, reducing residual chlorine with 0.008% sodium thiosulfate, storing in the dark, and adjusting the pH to 8-9; samples preserved in this manner may be held for seven days before extraction and for forty days after extraction. Exceptions to this optional preservation and holding time procedure are noted in footnote 5 (re the requirement for thiosulfate reduction of residual chlorine), and footnotes 12, 13 (re the analyses of benzene).

* If 1,2-diphenylhydrazine is likely to be present, adjust the pH of the sample to 4.0±0.2 to prevent rearrangement to benzidine.

* Extracts may be stored up to 7 days before analysis if storage is conducted under an inert (oxygen-free) atmosphere.

* For the analyses of diphenylhydrazine, add 0.008% Na₂S₂O₄ and adjust pH to 7-10 with NaOH within 24 hours of sampling.

* The pH adjustment may be performed upon receipt at the laboratory and may be omitted if the samples are extracted within 72 hours of collection. For the analyses of aldrin, add 0.008% Na₂S₂O₄.

8. Appendices A, B, and C are added to Part 136 to read as follows:

APPENDIX A TO PART 136—METHODS FOR ORGANIC CHEMICAL ANALYSIS OF MUNICIPAL AND INDUSTRIAL WASTEWATER.

Method 661—Purgeable Halocarbons

1. Scope and Application

1.1 This method covers the determination of 29 purgeable halocarbons.

The following parameters may be determined by this method:

Parameter	STORET No.	CAS No.
Bromochloromethane	32101	75-27-4
Bromodichloromethane	32104	75-25-2
Bromotrichloromethane	34413	74-83-9
Carbon tetrachloride	32102	56-23-5
Chlorobenzene	34301	108-90-7
Chloroethane	34311	75-00-3
2-Chloroethyl ether	34578	100-75-8
Chloroform	32106	67-66-3
Chloromethane	34418	74-87-3
Dibromochloromethane	32105	124-48-1
1,2-Dichlorobenzene	34536	95-50-1
1,3-Dichlorobenzene	34586	541-73-1
1,4-Dichlorobenzene	34571	106-46-7
Dichlorodifluoromethane	34688	75-71-8
1,1-Dichloroethane	34498	75-34-3
1,2-Dichloroethane	34521	107-06-2
1,1-Dichloroethene	34501	75-35-4
trans-1,2-Dichloroethene	34546	156-60-5
1,2-Dichloropropane	34541	78-67-5
cis-1,2-Dichloropropane	34704	10061-01-5
trans-1,2-Dichloropropane	34686	10061-02-6
Methylene chloride	34422	75-08-2
1,1,2,2-Tetrachloroethane	34516	78-34-5
Tetrachloroethene	34475	127-18-4
1,1,1-Trichloroethane	34508	71-55-6
1,1,2-Trichloroethane	34511	78-00-5
Tetrachloroethane	38180	78-01-6
Tetrachlorofluoromethane	34488	75-88-4
Vinyl chloride	38715	75-01-4

1.2 This is a purge and trap gas chromatographic (GC) method applicable to the determination of the compounds listed above in municipal and industrial discharges as provided under 40 CFR 136.1. When this method is used to analyze unfamiliar samples for any or all of the compounds above, compound identifications should be supported by at least one additional qualitative technique. This method describes analytical conditions for a second gas chromatographic column that can be used to confirm measurements made with the

primary column. Method 624 provides gas chromatograph/mass spectrometer (GC/MS) conditions appropriate for the qualitative and quantitative confirmation of results for most of the parameters listed above.

1.3 The method detection limit (MDL, defined in Section 12.1) for each parameter is listed in Table 1. The MDL for a specific wastewater may differ from those listed, depending upon the nature of interferences in the sample matrix.

1.4 Any modification of this method, beyond those expressly permitted, shall be considered as a major modification subject to application and approval of alternate test procedures under 40 CFR 136.4 and 136.5.

1.5 This method is restricted to use by or under the supervision of analysts experienced in the operation of a purge and trap system and a gas chromatograph and in the interpretation of gas chromatograms. Each analyst must demonstrate the ability to generate acceptable results with this method using the procedure described in Section 8.2.

2. Summary of Method

2.1 An inert gas is bubbled through a 5-mL water sample contained in a specially-designed purging chamber at ambient temperature. The halocarbons are efficiently transferred from the aqueous phase to the vapor phase. The vapor is swept through a sorbent trap where the halocarbons are trapped. After purging is completed, the trap is heated and backflushed with the inert gas to desorb the halocarbons onto a gas chromatographic column. The gas chromatograph is temperature programmed to separate the halocarbons which are then detected with a halide-specific detector.^{2,3}

2.2 The method provides an optional gas chromatographic column that may be helpful in resolving the compounds of interest from interferences that may occur.

3. Interferences

3.1 Impurities in the purge gas and organic compounds outgassing from the plumbing ahead of the trap account for the majority of contamination problems. The analytical system must be demonstrated to be free from contamination under the conditions of the analysis by running laboratory reagent blanks as described in Section 8.1.3. The use of non-Teflon plastic

tubing, non-Teflon thread sealants, or flow controllers with rubber components in the purge and trap system should be avoided.

3.2 Samples can be contaminated by diffusion of volatile organics (particularly fluorocarbons and methylene chloride) through the septum seal into the sample during shipment and storage. A field reagent blank prepared from reagent water and carried through the sampling and handling protocol can serve as a check on such contamination.

3.3 Contamination by carry-over can occur whenever high level and low level samples are sequentially analyzed. To reduce carry-over, the purging device and sample syringe must be rinsed with reagent water between sample analyses. Whenever an unusually concentrated sample is encountered, it should be followed by an analysis of reagent water to check for cross contamination. For samples containing large amounts of water-soluble materials, suspended solids, high boiling compounds or high organohalide levels, it may be necessary to wash out the purging device with a detergent solution, rinse it with distilled water, and then dry it in a 105°C oven between analyses. The trap and other parts of the system are also subject to contamination; therefore, frequent bakeout and purging of the entire system may be required.

4. Safety

4.1 The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical compound should be treated as a potential health hazard. From this viewpoint, exposure to these chemicals must be reduced to the lowest possible level by whatever means available. The laboratory is responsible for maintaining a current awareness file of OSHA regulations regarding the safe handling of the chemicals specified in this method. A reference file of material data handling sheets should also be made available to all personnel involved in the chemical analysis. Additional references to laboratory safety are available and have been identified^{4,5} for the information of the analyst.

4.2 The following parameters covered by this method have been tentatively classified as known or suspected, human or mammalian carcinogens: carbon tetrachloride, chloroform, 1,4-dichlorobenzene, and vinyl chloride. Primary standards of these toxic compounds should be prepared in a hood. A NIOSH/MESA approved toxic gas respirator should be worn when the analyst handles high concentrations of these toxic compounds.

5. Apparatus and Materials

5.1 Sampling equipment, for discrete sampling.

5.1.1 Vial—25-mL capacity or larger, equipped with a screw cap with a hole in the center (Pierce #13075 or equivalent). Detergent wash, rinse with tap and distilled water, and dry at 105 °C before use.

5.1.2 Septum—Teflon-faced silicone (Pierce #12722 or equivalent). Detergent wash, rinse with tap and distilled water, and dry at 105 °C for 1 h before use.

5.2 Purge and trap system—The purge and trap system consists of three separate pieces of equipment: a purging device, trap, and desorber. Several complete systems are now commercially available.

5.2.1 The purging device must be designed to accept 5-mL samples with a water column at least 3 cm deep. The gaseous head space between the water column and the trap must have a total volume of less than 15 mL. The purge gas must pass through the water column as finely divided bubbles with a diameter of less than 3 mm at the origin. The purge gas must be introduced no more than 5 mm from the base of the water column. The purging device illustrated in Figure 1 meets these design criteria.

5.2.2 The trap must be at least 25 cm long and have an inside diameter of at least 0.105 in. The trap must be packed to contain the following minimum lengths of adsorbents: 1.0 cm of methyl silicone coated packing (Section 6.3.3), 7.7 cm of 2,6-diphenylene oxide polymer (Section 6.3.2), 7.7 cm of silica gel (Section 6.3.4), 7.7 cm of coconut charcoal (Section 6.3.1). If it is not necessary to analyze for dichlorodifluoromethane, the charcoal can be eliminated, and the polymer section lengthened to 15 cm. The minimum specifications for the trap are illustrated in Figure 2.

5.2.3 The desorber must be capable of rapidly heating the trap to 180 °C. The polymer section of the trap should not be heated higher than 180 °C and the remaining sections should not exceed 200 °C. The desorber illustrated in Figure 2 meets these design criteria.

5.2.4 The purge and trap system may be assembled as a separate unit or be coupled to a gas chromatograph as illustrated in Figures 3 and 4.

5.3 Gas chromatograph—An analytical system complete with a temperature programmable gas chromatograph suitable for on-column injection and all required accessories including syringes, analytical columns, gases, detector, and strip-chart recorder. A data system is recommended for measuring peak areas.

5.3.1 Column 1—8 ft long x 0.1 in. ID stainless steel or glass, packed with 1% SP-100 on Carbowax B (60/80 mesh) or

equivalent. This column was used to develop the method performance statements in Section 12. Guidelines for the use of alternate column packings are provided in Section 10.1.

5.3.2 Column 2—8 ft long x 0.1 in. ID stainless steel or glass, packed with chemically bonded n-octane on Porasil-C (100/120 mesh) or equivalent.

5.3.3 Detector—Electrolytic conductivity or microcoulometric detector. These types of detectors have proven effective in the analysis of wastewaters for the parameters listed in the scope (Section 1.1). The electrolytic conductivity detector was used to develop the method performance statements in Section 12. Guidelines for the use of alternate detectors are provided in Section 10.1.

5.4 Syringes—5-mL glass hypodermic with Luerlok tip (two each), if applicable to the purging device.

5.5 Micro syringes—25-μL, 0.006 in. ID needle.

5.6 Syringe valve—2-way, with Luer ends (three each).

5.7 Syringe—5-mL, gas-tight with shut-off valve.

5.8 Bottle—15-mL, screw-cap, with Teflon cap liner.

5.9 Balance—Analytical, capable of accurately weighing 0.0001 g.

6. Reagents

6.1 Reagent water—Reagent water is defined as a water in which an interferent is not observed at the MDL of the parameters of interest.

6.1.1 Reagent water can be generated by passing tap water through a carbon filter bed containing about 1 lb of activated carbon (Filtrosorb-300, Calgon Corp., or equivalent).

6.1.2 A water purification system (Millipore Super-Q or equivalent) may be used to generate reagent water.

6.1.3 Reagent water may also be prepared by boiling water for 15 min. Subsequently, while maintaining the temperature at 90 °C, bubble a contaminant-free inert gas through the water for 1 h. While still hot, transfer the water to a narrow mouth screw-cap bottle and seal with a Teflon-lined septum and cap.

6.2 Sodium thiosulfate—(ACS) Granular.

6.3 Trap Materials:

6.3.1 Coconut charcoal—6/10 mesh sieved to 28 mesh, Barnebey Cheney, CA-580-26 lot # M-2649 or equivalent.

6.3.2 2,6-Diphenylene oxide polymer—Tenax, (60/80 mesh), chromatographic grade or equivalent.

6.3.3 Methyl silicone packing—3% OV-1 on Chromosorb-W (60/80 mesh) or equivalent.

6.3.4 Silica gel—35/60 mesh, Davison, grade-15 or equivalent.

6.4 Methanol—Pesticide quality or equivalent.

6.5 Stock standard solutions—Stock standard solutions may be prepared from pure standard materials or purchased as certified solutions. Prepare stock standard solutions in methanol using assayed liquids or gases as appropriate. Because of the toxicity of some of the organohalides, primary dilutions of these materials should be prepared in a hood. A NIOSH/MESA approved toxic gas respirator should be used

when the analyst handles high concentrations of such materials.

6.5.1 Place about 0.8 mL of methanol into a 10 mL ground glass stoppered volumetric flask. Allow the flask to stand, unstoppered, for about 10 min or until all alcohol wetted surfaces have dried. Weigh the flask to the nearest 0.1 mg.

6.5.2 Add the assayed reference material:

6.5.2.1 Liquid—Using a 100 μL syringe, immediately add two or more drops of assayed reference material to the flask, then reweigh. Be sure that the drops fall directly into the alcohol without contacting the neck of the flask.

6.5.2.2 Gases—To prepare standards for any of the six halocarbons that boil below 30 °C (bromomethane, chloromethane, chloromethane, dichlorodifluoromethane, trichlorofluoromethane, vinyl chloride), fill a 5-mL valved gas-tight syringe with the reference standard to the 5.0-mL mark. Lower the needle to 5 mm above the methanol meniscus. Slowly introduce the reference standard above the surface of the liquid (the heavy gas will rapidly dissolve into the methanol).

6.5.3 Reweigh, dilute to volume, stopper, then mix by inverting the flask several times. Calculate the concentration in μg/μL from the net gain in weight. When compound purity is assayed to be 96% or greater, the weight can be used without correction to calculate the concentration of the stock standard. Commercially prepared stock standards can be used at any concentration if they are certified by the manufacturer or by an independent source.

6.5.4 Transfer the stock standard solution into a Teflon-sealed screw-cap bottle. Store, with minimal headspace, at -10 to -20 °C and protect from light.

6.5.5 Prepare fresh standards weekly for the six gases and 2-chloroethylvinyl ether. All other standards must be replaced after one month, or sooner if comparison with check standards indicates a problem.

6.6 Secondary dilution standards—Using stock standard solutions, prepare secondary dilution standards in methanol that contain the compounds of interest, either singly or mixed together. The secondary dilution standards should be prepared at concentrations such that the aqueous calibration standards prepared in Sections 7.3.1 or 7.4.1 will bracket the working range of the analytical system. Secondary dilution standards should be stored with minimal headspace and should be checked frequently for signs of degradation or evaporation, especially just prior to preparing calibration standards from them.

6.7 Quality control check sample concentrate—See Section 8.2.1.

7. Calibration

7.1 Assemble a purge and trap system that meets the specifications in Section 5.2. Condition the trap overnight at 180 °C by backflushing with an inert gas flow of at least 20 mL/min. Condition the trap for 10 min once daily prior to use.

7.2 Connect the purge and trap system to a gas chromatograph. The gas chromatograph must be operated using temperature and flow

rate conditions equivalent to those given in Table 1. Calibrate the purge and trap-gas chromatographic system using either the external standard technique (Section 7.3) or the internal standard technique (Section 7.4).

7.3 External standard calibration procedure:

7.3.1 Prepare calibration standards at a minimum of three concentration levels for each parameter by carefully adding 20.0 μ L of one or more secondary dilution standards to 100, 500, or 1000 mL of reagent water. A 25- μ L syringe with a 0.006 in. ID needle should be used for this operation. One of the external standards should be at a concentration near, but above, the MDL (Table 1) and the other concentrations should correspond to the expected range of concentrations found in real samples or should define the working range of the detector. These aqueous standards can be stored up to 24 h, if held in sealed vials with zero headspace as described in Section 9.2. If not so stored, they must be discarded after 1 h.

7.3.2 Analyze each calibration standard according to Section 10, and tabulate peak height or area responses versus the concentration in the standard. The results can be used to prepare a calibration curve for each compound. Alternatively, if the ratio of response to concentration (calibration factor) is a constant over the working range ($<10\%$ relative standard deviation, RSD), linearity through the origin can be assumed and the average ratio or calibration factor can be used in place of a calibration curve.

7.4 Internal standard calibration procedure—To use this approach, the analyst must select one or more internal standards that are similar in analytical behavior to the compounds of interest. The analyst must further demonstrate that the measurement of the internal standard is not affected by method or matrix interferences. Because of these limitations, no internal standard can be suggested that is applicable to all samples. The compounds recommended for use as surrogate spikes in Section 8.7 have been used successfully as internal standards, because of their generally unique retention times.

7.4.1 Prepare calibration standards at a minimum of three concentration levels for each parameter of interest as described in Section 7.3.1.

7.4.2 Prepare a spiking solution containing each of the internal standards using the procedures described in Sections 6.5 and 6.6. It is recommended that the secondary dilution standard be prepared at a concentration of 15 μ g/mL of each internal standard compound. The addition of 10 μ L of this standard to 5.0 mL of sample or calibration standard would be equivalent to 30 μ g/L.

7.4.3 Analyze each calibration standard according to Section 10, adding 10 μ L of internal standard spiking solution directly to the syringe (Section 10.4). Tabulate peak height or area responses against concentration for each compound and internal standard, and calculate response factors (RF) for each compound using Equation 1.

Equation 1.

$$RF = \frac{(A_s)(C_u)}{(A_u)(C_s)}$$

where:

A_s = Response for the parameter to be measured.

A_u = Response for the internal standard.

C_u = Concentration of the internal standard.

C_s = Concentration of the parameter to be measured.

If the RF value over the working range is a constant ($<10\%$ RSD), the RF can be assumed to be invariant and the average RF can be used for calculations. Alternatively, the results can be used to plot a calibration curve of response ratios, A_s/A_u , vs. RF.

7.5 The working calibration curve, calibration factor, or RF must be verified on each working day by the measurement of a QC check sample.

7.5.1 Prepare the QC check sample as described in Section 8.2.2.

7.5.2 Analyze the QC check sample according to Section 10.

7.5.3 For each parameter, compare the response (Q) with the corresponding calibration acceptance criteria found in Table 2. If the responses for all parameters of interest fall within the designated ranges, analysis of actual samples can begin. If any individual Q falls outside the range, proceed according to Section 7.5.4.

Note: The large number of parameters in Table 2 present a substantial probability that one or more will not meet the calibration acceptance criteria when all parameters are analyzed.

7.5.4 Repeat the test only for those parameters that failed to meet the calibration acceptance criteria. If the response for a parameter does not fall within the range in this second test, a new calibration curve, calibration factor, or RF must be prepared for that parameter according to Section 7.3 or 7.4.

8. Quality Control

8.1 Each laboratory that uses this method is required to operate a formal quality control program. The minimum requirements of this program consist of an initial demonstration of laboratory capability and an ongoing analysis of spiked samples to evaluate and document data quality. The laboratory must maintain records to document the quality of data that is generated. Ongoing data quality checks are compared with established performance criteria to determine if the results of analyses meet the performance characteristics of the method. When results of sample spikes indicate atypical method performance, a quality control check standard must be analyzed to confirm that the measurements were performed in an in-control mode of operation.

8.1.1 The analyst must make an initial, one-time, demonstration of the ability to generate acceptable accuracy and precision with this method. This ability is established as described in Section 8.2.

8.1.2 In recognition of advances that are occurring in chromatography, the analyst is permitted certain options (detailed in Section 10.1) to improve the separations or lower the cost of measurements. Each time such a

modification is made to the method, the analyst is required to repeat the procedure in Section 8.2.

8.1.3 Each day, the analyst must analyze a reagent water blank to demonstrate that interferences from the analytical system are under control.

8.1.4 The laboratory must, on an ongoing basis, spike and analyze a minimum of 10% of all samples to monitor and evaluate laboratory data quality. This procedure is described in Section 8.3.

8.1.5 The laboratory must, on an ongoing basis, demonstrate through the analyses of quality control check standards that the operation of the measurement system is in control. This procedure is described in Section 8.4. The frequency of the check standard analyses is equivalent to 10% of all samples analyzed but may be reduced if spike recoveries from samples (Section 8.3) meet all specified quality control criteria.

8.1.6 The laboratory must maintain performance records to document the quality of data that is generated. This procedure is described in Section 8.5.

8.2 To establish the ability to generate acceptable accuracy and precision, the analyst must perform the following operations.

8.2.1 A quality control (QC) check sample concentrate is required containing each parameter of interest at a concentration of 10 μ g/mL in methanol. The QC check sample concentrate must be obtained from the U.S. Environmental Protection Agency, Environmental Monitoring and Support Laboratory in Cincinnati, Ohio, if available. If not available from that source, the QC check sample concentrate must be obtained from another external source. If not available from either source above, the QC check sample concentrate must be prepared by the laboratory using stock standards prepared independently from those used for calibration.

8.2.2 Prepare a QC check sample to contain 20 μ g/L of each parameter by adding 200 μ L of QC check sample concentrate to 100 mL of reagent water.

8.2.3 Analyze four 5-mL aliquots of the well-mixed QC check sample according to Section 10.

8.2.4 Calculate the average recovery (\bar{X}) in μ g/L and the standard deviation of the recovery (s) in μ g/L for each parameter of interest using the four results.

8.2.5 For each parameter compare s and \bar{X} with the corresponding acceptance criteria for precision and accuracy, respectively, found in Table 2. If s and \bar{X} for all parameters of interest meet the acceptance criteria, the system performance is acceptable and analysis of actual samples can begin. If any individual s exceeds the precision limit or any individual \bar{X} falls outside the range for accuracy, then the system performance is unacceptable for that parameter.

Note: The large number of parameters in Table 2 present a substantial probability that one or more will fail at least one of the acceptance criteria when all parameters are analyzed.

8.2.6 When one or more of the parameters tested fail at least one of the acceptance

criteria, the analyst must proceed according to Section 8.2.6.1 or 8.2.6.2.

8.2.6.1 Locate and correct the source of the problem and repeat the test for all parameters of interest beginning with Section 8.2.3.

8.2.6.2 Beginning with Section 8.2.3, repeat the test only for those parameters that failed to meet criteria. Repeated failure, however, will confirm a general problem with the measurement system. If this occurs, locate and correct the source of the problem and repeat the test for all compounds of interest beginning with Section 8.2.3.

8.3 The laboratory must, on an ongoing basis, spike at least 10% of the samples from each sample site being monitored to assess accuracy. For laboratories analyzing one to ten samples per month, at least one spiked sample per month is required.

8.3.1 The concentration of the spike in the sample should be determined as follows:

8.3.1.1 If, as in compliance monitoring, the concentration of a specific parameter in the sample is being checked against a regulatory concentration limit, the spike should be at that limit or 1 to 5 times higher than the background concentration determined in Section 8.3.2, whichever concentration would be larger.

8.3.1.2 If the concentration of a specific parameter in the sample is not being checked against a limit specific to that parameter, the spike should be at 20 µg/L or 1 to 5 times higher than the background concentration determined in Section 8.3.2, whichever concentration would be larger.

8.3.2 Analyze one 5-mL sample aliquot to determine the background concentration (B) of each parameter. If necessary, prepare a new QC check sample concentrate (Section 8.2.1) appropriate for the background concentrations in the sample. Spike a second 5-mL sample aliquot with 10 µL of the QC check sample concentrate and analyze it to determine the concentration after spiking (A) of each parameter. Calculate each percent recovery (P) as $100(A-B)/T$, where T is the known true value of the spike.

8.3.3 Compare the percent recovery (P) for each parameter with the corresponding QC acceptance criteria found in Table 2. These acceptance criteria were calculated to include an allowance for error in measurement of both the background and spike concentrations, assuming a spike to background ratio of 5:1. This error will be accounted for to the extent that the analyst's spike to background ratio approaches 5:1.⁷ If spiking was performed at a concentration lower than 20 µg/L, the analyst must use either the QC acceptance criteria in Table 2, or optional QC acceptance criteria calculated for the specific spike concentration. To calculate optional acceptance criteria for the recovery of a parameter: (1) Calculate accuracy (X') using the equation in Table 3, substituting the spike concentration (T) for C; (2) calculate overall precision (S') using the equation in Table 3, substituting X' for X; (3) calculate the range for recovery at the spike concentration as $(100 X' / T) \pm 2.44(100 S' / T)\%$.

8.3.4 If any individual P falls outside the designated range for recovery, that parameter has failed the acceptance criteria. A check

standard containing each parameter that failed the criteria must be analyzed as described in Section 8.4.

8.4 If any parameter fails the acceptance criteria for recovery in Section 8.3, a QC check standard containing each parameter that failed must be prepared and analyzed.

Note: The frequency for the required analysis of a QC check standard will depend upon the number of parameters being simultaneously tested, the complexity of the sample matrix, and the performance of the laboratory. If the entire list of parameters in Table 2 must be measured in the sample in Section 8.3, the probability that the analysis of a QC check standard will be required is high. In this case the QC check standard should be routinely analyzed with the spiked sample.

8.4.1 Prepare the QC check standard by adding 10 µL of QC check sample concentrate (Sections 8.2.1 or 8.3.2) to 5 mL of reagent water. The QC check standard needs only to contain the parameters that failed criteria in the test in Section 8.3.

8.4.2 Analyze the QC check standard to determine the concentration measured (A) of each parameter. Calculate each percent recovery (P_c) as $100(A/T)\%$, where T is the true value of the standard concentration.

8.4.3 Compare the percent recovery (P_c) for each parameter with the corresponding QC acceptance criteria found in Table 2. Only parameters that failed the test in Section 8.3 need to be compared with these criteria. If the recovery of any such parameter falls outside the designated range, the laboratory performance for that parameter is judged to be out of control, and the problem must be immediately identified and corrected. The analytical result for that parameter in the unspiked sample is suspect and may not be reported for regulatory compliance purposes.

8.5 As part of the QC program for the laboratory, method accuracy for wastewater samples must be assessed and records must be maintained. After the analysis of five spiked wastewater samples as in Section 8.3, calculate the average percent recovery (P̄) and the standard deviation of the percent recovery (s_p). Express the accuracy assessment as a percent recovery interval from $P̄ - 2s_p$ to $P̄ + 2s_p$. If $p = 90\%$ and $s_p = 10\%$, for example, the accuracy interval is expressed as 70–110%. Update the accuracy assessment for each parameter on a regular basis (e.g. after each five to ten new accuracy measurements).

8.6 It is recommended that the laboratory adopt additional quality assurance practices for use with this method. The specific practices that are most productive depend upon the needs of the laboratory and the nature of the samples. Field duplicates may be analyzed to assess the precision of the environmental measurements. When doubt exists over the identification of a peak on the chromatogram, confirmatory techniques such as gas chromatography with a dissimilar column, specific element detector, or mass spectrometer must be used. Whenever possible, the laboratory should analyze standard reference materials and participate in relevant performance evaluation studies.

8.7 The analyst should monitor both the performance of the analytical system and the

effectiveness of the method in dealing with each sample matrix by spiking each sample, standard, and reagent water blank with surrogate halocarbons. A combination of bromochloromethane, 2-bromo-1-chloropropane, and 1,4-dichlorobutane is recommended to encompass the range of the temperature program used in this method. From stock standard solutions prepared as in Section 6.5, add a volume to give 750 µg of each surrogate to 45 mL of reagent water contained in a 50-mL volumetric flask, mix and dilute to volume for a concentration of 15 ng/µL. Add 10 µL of this surrogate spiking solution directly into the 5-mL syringe with every sample and reference standard analyzed. Prepare a fresh surrogate spiking solution on a weekly basis. If the internal standard calibration procedure is being used, the surrogate compounds may be added directly to the internal standard spiking solution (Section 7.4.2).

9. Sample Collection, Preservation, and Handling

9.1 All samples must be iced or refrigerated from the time of collection until analysis. If the sample contains free or combined chlorine, add sodium thiosulfate preservative (10 mg/40 mL is sufficient for up to 5 ppm Cl₂) to the empty sample bottle just prior to shipping to the sampling site. EPA Methods 330.4 and 330.5 may be used for measurement of residual chlorine.⁸ Field test kits are available for this purpose.

9.2 Grab samples must be collected in glass containers having a total volume of at least 25 mL. Fill the sample bottle just to overflowing in such a manner that no air bubbles pass through the sample as the bottle is being filled. Seal the bottle so that no air bubbles are entrapped in it. If preservative has been added, shake vigorously for 1 min. Maintain the hermetic seal on the sample bottle until time of analysis.

9.3 All samples must be analyzed within 14 days of collection.⁹

10. Procedure

10.1 Table 1 summarizes the recommended operating conditions for the gas chromatograph. Included in this table are estimated retention times and MDL that can be achieved under these conditions. An example of the separations achieved by Column 1 is shown in Figure 5. Other packed columns, chromatographic conditions, or detectors may be used if the requirements of Section 8.2 are met.

10.2 Calibrate the system daily as described in Section 7.

10.3 Adjust the purge gas (nitrogen or helium) flow rate to 40 mL/min. Attach the trap inlet to the purging device, and set the purge and trap system to purge (Figure 3). Open the syringe valve located on the purging device sample introduction needle.

10.4 Allow the sample to come to ambient temperature prior to introducing it to the syringe. Remove the plunger from a 5-mL syringe and attach a closed syringe valve. Open the sample bottle (or standard) and carefully pour the sample into the syringe barrel to just short of overflowing. Replace the syringe plunger and compress the sample. Open the syringe valve and vent any residu-

air while adjusting the sample volume to 5.0 mL. Since this process of taking an aliquot destroys the validity of the sample for future analysis, the analyst should fill a second syringe at this time to protect against possible loss of data. Add 10.0 µL of the surrogate spiking solution (Section 8.7) and 10.0 µL of the internal standard spiking solution (Section 7.4.2), if applicable, through the valve bore, then close the valve.

10.5 Attach the syringe-syringe valve assembly to the syringe valve on the purging device. Open the syringe valves and inject the sample into the purging chamber.

10.6 Close both valves and purge the sample for 11.0 ± 0.1 min at ambient temperature.

10.7 After the 11-min purge time, attach the trap to the chromatograph, adjust the purge and trap system to the desorb mode (Figure 4), and begin to temperature program the gas chromatograph. Introduce the trapped materials to the GC column by rapidly heating the trap to 180 °C while backflushing the trap with an inert gas between 20 and 60 mL/min for 4 min. If rapid heating of the trap cannot be achieved, the GC column must be used as a secondary trap by cooling it to 30 °C (subambient temperature, if poor peak geometry or random retention time problems persist) instead of the initial program temperature of 45 °C.

10.8 While the trap is being desorbed into the gas chromatograph, empty the purging chamber using the sample introduction syringe. Wash the chamber with two 5-mL flushes of reagent water.

10.9 After desorbing the sample for 4 min, condition the trap by returning the purge and trap system to the purge mode. Wait 15 s then close the syringe valve on the purging device to begin gas flow through the trap. The trap temperature should be maintained at 180 °C. After approximately 7 min, turn off the trap heater and open the syringe valve to stop the gas flow through the trap. When the trap is cool, the next sample can be analyzed.

10.10 Identify the parameters in the sample by comparing the retention times of the peaks in the sample chromatogram with those of the peaks in standard chromatograms. The width of the retention time window used to make identifications should be based upon measurements of actual retention time variations of standards over the course of a day. Three times the standard deviation of a retention time for a compound can be used to calculate a suggested window size; however, the experience of the analyst should weigh heavily in the interpretation of chromatograms.

10.11 If the response for a peak exceeds the working range of the system, prepare a dilution of the sample with reagent water from the aliquot in the second syringe and reanalyze.

11. Calculations

11.1 Determine the concentration of individual compounds in the sample.

11.1.1 If the external standard calibration procedure is used, calculate the concentration of the parameter being measured from the peak response using the calibration curve or calibration factor determined in Section 7.3.2.

11.1.2 If the internal standard calibration procedure is used, calculate the concentration in the sample using the response factor (RF) determined in Section 7.4.3 and Equation 2.

Equation 2.

$$\text{Concentration } (\mu\text{g/L}) = \frac{(A_s)(C_{is})}{(A_{is})(RF)}$$

where:

A_s = Response for the parameter to be measured.

A_{is} = Response for the internal standard.

C_{is} = Concentration of the internal standard.

11.2 Report results in µg/L without correction for recovery data. All QC data obtained should be reported with the sample results.

12. Method Performance

12.1 The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero.¹ The MDL concentrations listed in Table 1 were obtained using reagent water.² Similar results were achieved using representative wastewaters. The MDL actually achieved in a given analysis will vary depending on instrument sensitivity and matrix effects.

12.2 This method is recommended for use in the concentration range from the MDL to 1000 × MDL. Direct aqueous injection techniques should be used to measure concentration levels above 1000 × MDL.

12.3 This method was tested by 20 laboratories using reagent water, drinking water, surface water, and three industrial wastewaters spiked at six concentrations over the range 8.0 to 500 µg/L.³ Single

operator precision, overall precision, and method accuracy were found to be directly related to the concentration of the parameter and essentially independent of the sample matrix. Linear equations to describe these relationships are presented in Table 3.

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TABLE 1.—CHROMATOGRAPHIC CONDITIONS AND METHOD DETECTION LIMITS

Parameter	Retention time (min)		Method detection limit (µg/L)
	Column 1	Column 2	
Chloromethane	1.30	5.28	0.08
Bromomethane	2.17	7.05	1.18
Dichlorodifluoromethane	2.62	nd	1.81
Vinyl chloride	2.67	5.28	0.18
Chloroethane	3.33	8.66	0.52
Methylene chloride	5.25	10.1	0.25
Trichlorofluoromethane	7.18	nd	nd
1,1-Dichloroethane	7.83	7.72	0.13
1,1-Dichloroethane	8.30	12.6	0.07
trans-1,2-Dichloroethane	10.1	9.38	0.10

TABLE 1.—CHROMATOGRAPHIC CONDITIONS AND METHOD DETECTION LIMITS—Continued

Parameter	Retention time (min)		Method detection limit (µg/L)
	Column 1	Column 2	
Chloroform	10.7	12.1	0.05
1,2-Dichloroethane	11.4	15.4	0.03
1,1,1-Trichloroethane	12.6	13.1	0.03
Carbon tetrachloride	13.0	14.4	0.12
Bromodichloromethane	13.7	14.6	0.10
1,2-Dichloropropane	14.8	16.6	0.04
cis-1,3-Dichloropropene	15.2	16.6	0.34
Trichloroethene	15.8	13.1	0.12
Dibromodichloromethane	16.5	16.8	0.08
1,1,2-Trichloroethane	16.5	18.1	0.02
trans-1,3-Dichloropropene	16.5	18.0	0.20
2-Chlorostyryl vinyl ether	18.0	nd	0.13
Bromotoluene	19.2	19.2	0.20
1,1,2,2-Tetrachloroethane	21.8	nd	0.03
Tetrachloroethane	21.7	15.0	0.03
Chlorobenzene	24.2	18.8	0.25
1,3-Dichlorobenzene	34.0	22.4	0.32
1,2-Dichlorobenzene	34.9	23.5	0.15
1,4-Dichlorobenzene	35.4	22.3	0.24

Column 1 conditions: Carbowax B (80/80 mesh) coated with 1% SP-1000 packed in an 8 ft x 0.1 in. ID stainless steel or glass column with helium carrier gas at 40 mL/min flow rate. Column temperature held at 45 °C for 3 min then programmed at 8 °C/min to 220 °C and held for 15 min.
 Column 2 conditions: Porasil-C (100/120 mesh) coated with n-octane packed in a 8 ft x 0.1 in. ID stainless steel or glass column with helium carrier gas at 40 mL/min flow rate. Column temperature held at 50 °C for 3 min then programmed at 8 °C/min to 170 °C and held for 4 min.
 nd = not determined.

TABLE 2.—CALIBRATION AND QC ACCEPTANCE CRITERIA—METHOD 601*

Parameter	Range for Q (µg/L)	Limit for s (µg/L)	Range for \bar{x} (µg/L)	Range for P _i (%)
Bromodichloromethane	15.2-24.8	4.3	10.7-32.0	42-172
Bromotoluene	14.7-25.3	4.7	5.0-29.3	13-158
Bromobenzene	11.7-26.3	7.6	3.4-24.5	D-144
Carbon tetrachloride	13.7-26.3	5.6	11.8-25.3	43-143
Chlorobenzene	14.4-25.6	5.0	10.2-27.4	38-150
Chloroethane	15.4-24.6	4.4	11.3-25.2	46-137
2-Chlorostyryl vinyl ether	12.0-28.0	8.3	4.5-35.5	14-188
Chloroform	15.0-25.0	4.5	12.4-24.0	48-133
Chloromethane	11.9-26.1	7.4	D-34.8	D-185
Dibromodichloromethane	13.1-26.8	6.3	7.9-35.1	24-191
1,2-Dichlorobenzene	14.0-26.0	5.5	1.7-38.9	D-208
1,3-Dichlorobenzene	9.8-30.1	9.1	8.2-32.6	7-187
1,4-Dichlorobenzene	13.9-26.1	5.5	11.5-25.5	42-143
1,1-Dichloroethane	16.8-23.2	3.2	11.2-24.8	47-132
1,2-Dichloroethane	14.3-25.7	5.2	13.0-28.5	51-147
1,1-Dichloroethene	12.6-27.4	6.6	10.2-27.3	28-167
trans-1,2-Dichloroethene	12.8-27.2	6.4	11.4-27.1	38-156
1,2-Dichloropropane	14.8-25.2	5.2	10.1-29.9	44-158
cis-1,3-Dichloropropene	12.8-27.2	7.3	6.2-33.8	22-178
trans-1,3-Dichloropropene	12.8-27.2	7.3	6.2-33.8	22-178
Methylene chloride	15.5-24.5	4.0	7.0-27.6	25-182
1,1,2,2-Tetrachloroethane	9.8-30.2	9.2	6.8-31.8	8-184
Tetrachloroethane	14.0-26.0	5.4	8.1-29.6	26-182
1,1,1-Trichloroethane	14.2-25.8	4.9	10.8-24.8	41-138
1,1,2-Trichloroethane	15.7-24.3	3.8	8.6-25.4	38-138
Trichloroethene	15.4-24.6	4.2	9.2-26.6	35-146
Trichlorodichloromethane	13.3-26.7	6.0	7.4-28.1	21-156
Vinyl chloride	13.7-26.3	5.7	6.2-29.9	28-183

Q = Concentration measured in QC check sample, in µg/L (Section 7.5.3).
 s = Standard deviation of four recovery measurements, in µg/L (Section 8.2.4).
 \bar{x} = Average recovery for four recovery measurements, in µg/L (Section 8.2.4).
 P_i = Percent recovery measured (Section 8.3.2, Section 8.4.2).
 D = Detected, result must be greater than zero.
 * Criteria were calculated assuming a QC check sample concentration of 20 µg/L.

Note: These criteria are based directly upon the method performance data in Table

3. Where necessary, the limits for recovery have been broadened to assure applicability

of the limits to concentrations below those used to develop Table 3.

TABLE 3.—METHOD ACCURACY AND PRECISION AS FUNCTIONS OF CONCENTRATION—METHOD 601

Parameter	Accuracy, as recovery, \bar{x} (µg/L)	Single analyst precision, s _i (µg/L)	Overall precision, S (µg/L)
Bromodichloromethane	1.12C - 1.02	0.11 \bar{x} + 0.04	0.20 \bar{x} + 1.00
Bromotoluene	0.96C - 2.05	0.12 \bar{x} + 0.58	0.21 \bar{x} + 2.41
Bromobenzene	0.78C - 1.27	0.26 \bar{x} + 0.27	0.36 \bar{x} + 0.94
Carbon tetrachloride	0.98C - 1.04	0.15 \bar{x} + 0.38	0.20 \bar{x} + 0.38
Chlorobenzene	1.00C - 1.23	0.15 \bar{x} - 0.02	0.18 \bar{x} + 1.21
Chloroethane	0.89C - 1.53	0.14 \bar{x} - 0.13	0.17 \bar{x} + 0.83
2-Chlorostyryl vinyl ether*	1.00C	0.20 \bar{x}	0.35 \bar{x}
Chloroform	0.93C - 0.39	0.13 \bar{x} + 0.15	0.18 \bar{x} - 0.02
Chloromethane	0.77C - 0.18	0.28 \bar{x} - 0.31	0.52 \bar{x} + 1.31
Dibromodichloromethane	0.94C - 2.72	0.11 \bar{x} + 1.10	0.24 \bar{x} + 1.68
1,2-Dichlorobenzene	0.83C - 1.70	0.20 \bar{x} - 0.97	0.13 \bar{x} + 6.13

TABLE 3.—METHOD ACCURACY AND PRECISION AS FUNCTIONS OF CONCENTRATION—METHOD 601—Continued

Parameter	Accuracy, as recovery, \bar{X} (%)	Single analyst precision, s_x (%)	Overall precision, S (%)
1,3-Dichlorobenzene	0.94C ± 0.43	0.14X ± 2.33	0.26X ± 2.34
1,4-Dichlorobenzene	0.93C ± 0.08	0.15X ± 0.28	0.20X ± 0.41
1,1-Dichloroethene	0.94C ± 1.08	0.08X ± 0.17	0.14X ± 0.84
1,2-Dichloroethene	1.04C ± 1.08	0.11X ± 0.70	0.15X ± 0.84
1,1-Dichloroethane	0.98C ± 0.87	0.21X ± 0.23	0.29X ± 0.40
trans-1,2-Dichloroethane	0.87C ± 0.16	0.11X ± 1.46	0.17X ± 1.46
1,2-Dichloropropane*	1.00C	0.13X	0.23X
cis-1,3-Dichloropropane*	1.00C	0.18X	0.32X
trans-1,3-Dichloropropane*	1.00C	0.18X	0.32X
Methylene chloride	0.91C ± 0.93	0.11X ± 0.33	0.21X ± 1.43
1,1,2,2-Tetrachloroethane	0.95C ± 0.19	0.14X ± 2.41	0.23X ± 2.79
Tetrachloroethene	0.94C ± 0.06	0.14X ± 0.38	0.18X ± 2.21
1,1,1-Trichloroethane	0.90C ± 0.16	0.15X ± 0.04	0.20X ± 0.37
1,1,2-Trichloroethane	0.88C ± 0.30	0.13X ± 0.14	0.19X ± 0.87
Trichloroethene	0.87C ± 0.48	0.13X ± 0.03	0.23X ± 0.30
Trichlorofluoromethane	0.89C ± 0.07	0.15X ± 0.67	0.26X ± 0.91
Vinyl chloride	0.87C ± 0.36	0.13X ± 0.65	0.27X ± 0.40

\bar{X} = Expected recovery for one or more measurements of a sample containing a concentration of C, in $\mu\text{g/L}$.

s_x = Expected single analyst standard deviation of measurements at an average concentration found of \bar{X} , in $\mu\text{g/L}$.

S = Expected interlaboratory standard deviation of measurements at an average concentration found of \bar{X} , in $\mu\text{g/L}$.

C = True value for the concentration, in $\mu\text{g/L}$.

X = Average recovery found for measurements of samples containing a concentration of C, in $\mu\text{g/L}$.

* Estimates based upon the performance in a single laboratory.¹⁰

